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Prio	or Applica	tion:	TYPED NAME HAMMID SANCHEZ
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or <u>l</u>	NOVEL METH	ODS OF DIAGNOSING AND TRE	ATING BREAST CANCER, COMPOSITIONS, AND
METH(ODS OF SCR	REENING FOR BREAST CANCER	MODULATORS.
1.	(a) <u>X</u>	Enclosed is a new appli	cation.
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3.	(a)	Enclosed is a Small Ent	ity Affidavit.
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Claims as filed in the prior application, less any claims canceled by amendment below:

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INDEP CLAIMS	3 =	*	x 39 =	\$	OR	x 78 =	\$
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Cancel in this application original claims ______ of the prior application before calculating the filing fee. (At least one independent claim must be retained for filing purposes.)

Amend the specification by inserting before the first line the sentence:

of pending applications Serial No. 09/268,865, filed March 15, 1999; Serial No. 09/439,878, filed November 12, 1999; Serial No. 09/440,370, filed November 12, 1999; Serial No. 09/440,493, filed November 15, 1999; Serial No. 09/440,676, filed November 16, 1999; Serial No. 09/440,677, filed November 16, 1999; Serial No. 09/450,810, filed November 29, 1999; Serial No. 09/453,137, filed December 2, 1999; and Serial No. NOT ASSIGNED, filed March 8, 2000.--

- 9. (a) X 188 sheets of drawings are enclosed.
- 10. (a) _____ filed on ___ in ____ is claimed under 35 U.S.C. 119.
 - (b) ____ The certified copy has been filed in prior application Serial No. ____ filed on ____.
- 11. ____ An Assignment is enclosed.

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12.		The prior application is assigned of record to				
13.		A Power of Attorney by Assignee is enclosed.				
14.		The power of attorney in the prior application is to:				
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	(a)	The power appears in the original papers in the prior application.				
	(b)	Since the power does not appear in the original papers, a copy of the power in the prior application is enclosed.				
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		FLEHR HOHBACH TEST ALBRITTON & HERBERT LLP Suite 3400, Four Embarcadero Center San Francisco, California 94111-4187 Telephone: (415) 781-1989				
5.		A preliminary amendment is enclosed. (Claims added by this amendment have been properly numbered consecutively beginning with the number next following the highest numbered original claim in the prior application.)				
16.		A Prior Art Statement is enclosed.				
17.		I hereby verify that the attached papers are a true duplicate of prior application Serial No as originally filed on				
Date	: <u>March</u>	Dolly A. Vance Registration No. 39.054				

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NOVEL METHODS OF DIAGNOSING AND TREATING BREAST CANCER, COMPOSITIONS, AND METHODS OF SCREENING FOR BREAST CANCER MODULATORS

FIELD OF THE INVENTION

The invention relates to the identification of expression profiles and the nucleic acids involved in breast cancer, and to the use of such expression profiles and nucleic acids in diagnosis, prognosis and treatment of breast cancer. The invention further relates to methods for identifying and using candidate agents and/or targets which modulate breast cancer.

BACKGROUND OF THE INVENTION

Breast cancer is a significant cancer in Western populations. It develops as the result of a pathologic transformation of normal breast epithelium to an invasive cancer. There have been a number of recently characterized genetic alterations that have been implicated in breast cancer. However, there is a need to identify all of the genetic alterations involved in the development of breast cancer.

Imaging of breast cancer for diagnosis has been problematic and limited. In addition, dissemination of tumor cells (metastases) to locoregional lymph nodes is an important prognostic factor; five year survival rates drop from 80 percent in patients with no lymph node metastases to 45 to 50 percent in those patients who do have lymph node metastases. A recent report showed that micrometastases can be detected from lymph nodes using reverse transcriptase-PCR methods based on the presence of mRNA for carcinoembryonic antigen, which has previously been shown to be present in the vast majority of breast cancers but not in normal tissues. Liefers et al., New England J. of Med. 339(4):223 (1998).

Thus, methods that can be used for diagnosis and prognosis of breast cancer would be desirable. While academia and industry has made an effort to identify novel sequences, there has not been an equal effort exerted to identify the function of the novel sequences. For example, databases show the

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sequence of accession number W72838, but there no data correlating this sequence with a function, much less a disease state. Accordingly, provided herein are methods that can be used in diagnosis and prognosis of breast cancer. Further provided are methods that can be used to screen candidate bioactive agents for the ability to modulate breast cancer. Additionally, provided herein are molecular targets for therapeutic intervention in breast and other cancers.

SUMMARY OF THE INVENTION

The present invention provides methods for screening for compositions which modulate breast cancer. Also provided herein are methods of inhibiting proliferation of a cell, preferably a breast cancer cell. Methods of treatment of cancer, as well as compositions, are also provided herein.

In one aspect, a method of screening drug candidates comprises providing a cell that expresses an expression profile gene or fragments thereof. Preferred embodiments of the expression profile gene are genes which are differentially expressed in cancer cells, preferably breast cancer cells, compared to other cells. Preferred embodiments of expression profile genes used in the methods herein include but are not limited to the group consisting of BCR3, BCQ8, BCQ5, BCH1, BCN1, BCN2, BCN5, BCO2, BCX2, BCJ7, BCY3, BCX3, BCA2 and BCR2; fragments of the proteins of this group are also preferred. In another embodiment, a nucleic acid is selected from Figures 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, or 12. Preferred nucleic acids are in Figure 10, more preferably Figure 11, and most preferably in Figure 12. The method further includes adding a drug candidate to the cell and determining the effect of the drug candidate on the expression of the expression profile gene.

In one embodiment, the method of screening drug candidates includes comparing the level of expression in the absence of the drug candidate to the level of expression in the presence of the drug candidate, wherein the concentration of the drug candidate can vary when present, and wherein the comparison can occur after addition or removal of the drug candidate. In a preferred embodiment, the cell expresses at least two expression profile genes. The profile genes may show an increase or decrease.

Also provided herein is a method of screening for a bioactive agent capable of binding to a breast cancer modulator protein (BCMP), the method comprising combining the BCMP and a candidate bioactive agent, and determining the binding of the candidate agent to the BCMP. Preferably the BCMP is a protein or fragment thereof selected from the group consisting of BCR3, BCQ8, BCQ5,

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BCH1, BCN1, BCN2, BCN5, BCO2, BCX2, BCJ7, BCY3, BCX3, BCA2 and BCR2. In another embodiment, the protein is encoded by a nucleic acid selected from Figures 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, or 12. Preferred nucleic acids are in Figure 10, more preferably Figure 11, and most preferably in Figure 12.

Further provided herein is a method for screening for a bioactive agent capable of modulating the activity of a BCMP. In one embodiment, the method comprises combining the BCMP and a candidate bioactive agent, and determining the effect of the candidate agent on the bioactivity of the BCMP. Preferably the BCMP is a protein or fragment thereof selected from the group consisting of BCR3, BCQ8, BCQ5, BCH1, BCN1, BCN2, BCN5, BCO2, BCX2, BCJ7, BCY3, BCX3, BCA2 and BCR2. In another embodiment, the protein is encoded by a nucleic acid selected from Figures 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, or 12. Preferred nucleic acids are in Figure 10, more preferably Figure 11, and most preferably in Figure 12.

Also provided is a method of evaluating the effect of a candidate breast cancer drug comprising administering the drug to a transgenic animal expressing or over-expressing the BCMP, or an animal lacking the BCMP, for example as a result of a gene knockout.

Additionally, provided herein is a method of evaluating the effect of a candidate breast cancer drug comprising administering the drug to a patient and removing a cell sample from the patient. The expression profile of the cell is then determined. This method may further comprise comparing the expression profile to an expression profile of a healthy individual.

Moreover, provided herein is a biochip comprising a nucleic acid segment which encodes a breast cancer protein, preferably selected from the group consisting of BCR3, BCQ8, BCQ5, BCH1, BCN1, BCN2, BCN5, BCO2, BCX2, BCJ7, BCY3, BCX3, BCA2 and BCR2, or a fragment thereof, wherein the biochip comprises fewer than 1000 nucleic acid probes. Preferably at least two nucleic acid segments are included. In another embodiment, the nucleic acid selected from Figures 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, or 12. Preferred nucleic acids are in Figure 10, more preferably Figure 11, and most preferably in Figure 12.

Furthermore, a method of diagnosing a disorder associated with breast cancer is provided. The method comprises determining the expression of a gene which encodes a breast cancer protein preferably selected from the group consisting of BCR3, BCQ5, BCH1, BCN1, BCN2, BCN5,

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BCO2, BCX2, BCJ7, BCY3, BCX3, BCA2 and BCR2 or a fragment thereof in a first tissue type of a first individual, and comparing the distribution to the expression of the gene from a second normal tissue type from the first individual or a second unaffected individual. In another embodiment, the protein is encoded by a nucleic acid selected from Figures 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, or 12. Preferred nucleic acids are in Figure 10, more preferably Figure 11, and most preferably in Figure 12. A difference in the expression indicates that the first individual has a disorder associated with breast cancer.

In another aspect, the present invention provides an antibody which specifically binds to a breast cancer protein, preferably selected from the group consisting of BCR3, BCQ8, BCQ5, BCH1, BCN1, BCN2, BCN5, BCO2, BCX2, BCJ7, BCY3, BCX3, BCA2 and BCR2, or a fragment thereof. In another embodiment, the protein is encoded by a nucleic acid selected from Figures 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, or 12. Preferred nucleic acids are in Figure 10, more preferably Figure 11, and most preferably in Figure 12. In a preferred embodiment, the fragment of BCH1 is selected from BCH1p1 and BCH1p2. Other preferred fragments for the breast cancer proteins are shown in the figures. Preferably the antibody is a monoclonal antibody. The antibody can be a fragment of an antibody such as a single stranded antibody as further described herein, or can be conjugated to another molecule. In one embodiment, the antibody is a humanized antibody.

In one embodiment a method for screening for a bioactive agent capable of interfering with the binding of a breast cancer protein (BCMP) or a fragment thereof and an antibody which binds to said BCMP or fragment thereof. In a preferred embodiment, the method comprises combining a BCMP or fragment thereof, a candidate bioactive agent and an antibody which binds to said BCMP or fragment thereof. The method further includes determining the binding of said BCMP or fragment thereof and said antibody. Wherein there is a change in binding, an agent is identified as an interfering agent. The interfering agent can be an agonist or an antagonist. Preferably, the antibody as well as the agent inhibits breast cancer.

In a further aspect, a method for inhibiting breast cancer is provided. In one embodiment, the method comprises administering to a cell a composition comprising an antibody to a breast modulating protein, preferably selected from the group consisting of BCR3, BCQ8, BCQ5, BCH1, BCN1, BCN2, BCN5, BCO2, BCX2, BCJ7, BCY3, BCX3, BCA2 and BCR2, or a fragment thereof. In another embodiment, the protein is encoded by a nucleic acid selected from Figures 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, or 12. Preferred nucleic acids are in Figure 10, more preferably Figure 11, and most preferably in Figure 12.

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The method can be performed in vitro or in vivo, preferably in vivo to an individual. In a preferred embodiment the method of inhibiting breast cancer is provided to an individual with cancer. As described herein, methods of inhibiting breast cancer can be performed by administering an inhibitor of breast cancer protein activity, including an antisense molecules, and preferably small molecules.

Also provided herein are methods eliciting an immune response in an individual. In one embodiment a method provided herein comprises administering to an individual a composition comprising a breast modulating protein, preferably selected from the group consisting of BCR3, BCQ8, BCQ5, BCH1, BCN1, BCN2, BCN5, BCO2, BCX2, BCJ7, BCY3, BCX3, BCA2 and BCR2, or a fragment thereof. In another embodiment, the protein is encoded by a nucleic acid selected from Figures 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, or 12. Preferred nucleic acids are in Figure 10, more preferably Figure 11, and most preferably in Figure 12. In another aspect, said composition comprises a nucleic acid comprising a sequence encoding a breast modulating protein, preferably selected from the group consisting of BCR3, BCQ8, BCQ5, BCH1, BCN1, BCN2, BCN5, BCO2, BCX2, BCJ7, BCY3, BCX3, BCA2 and BCR2, or a fragment thereof. In another embodiment, the nucleic acid is selected from Figures 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, or 12. Preferred nucleic acids are in Figure 10, more preferably Figure 11, and most preferably in Figure 12.

Further provided herein are compositions capable of eliciting an immune response in an individual. In one embodiment, a composition provided herein comprises a breast modulating protein, preferably selected from the group consisting of BCR3, BCQ8, BCQ5, BCH1, BCN1, BCN2, BCN5, BCO2, BCX2, BCJ7, BCY3, BCX3, BCA2 and BCR2, or a fragment thereof, and a pharmaceutically acceptable carrier. In another embodiment, the protein is encoded by a nucleic acid selected from Figures 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, or 12. Preferred nucleic acids are in Figure 10, more preferably Figure 11, and most preferably in Figure 12. In another embodiment, said composition comprises a nucleic acid comprising a sequence encoding a breast modulating protein, preferably selected from the group consisting of BCR3, BCQ8, BCQ5, BCH1, BCN1, BCN2, BCN5, BCO2, BCX2, BCJ7, BCY3, BCX3, BCA2 and BCR2, or a fragment thereof, and a pharmaceutically acceptable carrier. In another embodiment, the nucleic acid is selected from Figures 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, or 12. Preferred nucleic acids are in Figure 10, more preferably Figure 11, and most preferably in Figure 12.

A method of neutralizing the effect of a breast cancer protein, preferably selected from the group consisting of BCR3, BCQ8, BCQ5, BCH1, BCN1, BCN2, BCN5, BCO2, BCX2, BCJ7, BCY3, BCX3, BCA2 and BCR2, or a fragment thereof, is also provided. Preferably, the method comprises

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contacting an agent specific for said protein with said protein in an amount sufficient to effect neutralization. In another embodiment, the protein is encoded by a nucleic acid selected from Figures 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, or 12. Preferred nucleic acids are in Figure 10, more preferably Figure 11, and most preferably in Figure 12.

In another aspect of the invention, a method of treating an individual for breast cancer is provided. In one embodiment, the method comprises administering to said individual an inhibitor of BCH1. In another embodiment, the individual is non-responsive to an anti-estrogen and is positive for estrogen receptor. Optionally, the method further comprises administering an anti-estrogen to said individual.

Also provided herein is method for determining the prognosis of an individual with breast cancer comprising determining the level of BCH1 in a sample, wherein a high level of BCH1 indicates a poor prognosis. Moreover, in yet another aspect of the invention, a method is provided for determining whether an individual with breast cancer will be non-responsive to anti-estrogen therapies comprising determining the level of BCH1 wherein a high level of BCH1 indicates that an individual will be non-responsive. Preferably, the methods herein are used to identify persons who are tamoxifen resistant.

Novel sequences are also provided herein. Other aspects of the invention will become apparent to the skilled artisan by the following description of the invention.

DETAILED DESCRIPTION OF THE FIGURES

Figure 1 provides the Accession numbers for genes, including expression sequence tags, (incorporated in their entirety here and throughout the application where Accession numbers are provided), downregulated in tumor tissue compared to normal breast tissue.

Figure 2 provides the Accession numbers for genes, including expression sequence tags, downregulated in tumor tissue compared to normal breast tissue.

Figure 3 provides the Accession numbers for genes, including expression sequence tags, downregulated in tumor tissue compared to normal breast tissue.

Figure 4 provides the Accession numbers for genes, including expression sequence tags, upregulated in tumor tissue compared to normal breast tissue.

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Figure 5 provides the Accession numbers for genes, including expression sequence tags, upregulated in tumor tissue compared to normal breast tissue.

Figure 6 provides the Accession numbers for genes, including expression sequence tags, upregulated in tumor tissue compared to normal breast tissue.

Figure 7 provides the Accession numbers for genes, including expression sequence tags, upregulated in tumor tissue compared to normal breast tissue. Open reading frames in the sequences have been characterized as having a signal sequence (SS), a transmembrane domain (TM) or other.

Figure 8 provides the Accession numbers for genes, including expression sequence tags, upregulated in tumor tissue compared to normal breast tissue. Specifically, one column shows the ratio of expression of the indicated gene in breast tumor tissue compared to other body tissues, and another column shows the ratio of expression of the indicated gene in breast tumor tissue compared to normal breast tissue.

Figure 9 depicts the Accession numbers for genes, including expression sequence tags, upregulated in tumor tissue compared to normal breast tissue.

Figure 10 depicts a preferred group of 1007 Accession numbers for genes, including expression sequence tags, upregulated in tumor tissue compared to normal breast tissue.

Figure 11 depicts a preferred group of 123 Accession numbers for genes, including expression sequence tags, upregulated in tumor tissue compared to normal breast tissue.

Figure 12 depicts a preferred group of 10 Accession numbers for genes, including expression sequence tags, upregulated in tumor tissue compared to normal breast tissue.

Figure 13 shows an embodiment of a nucleic acid (mRNA) which includes a sequence which encodes a breast cancer protein provided herein, BCR3. The start and stop codons are underlined.

Figure 14 shows an embodiment of an open reading frame of a nucleic acid encoding BCR3, wherein the start and stop codons are underlined.

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Figure 15 shows an embodiment of an amino acid sequence of BCR3. The signal peptide is underlined and the transmembrane domain is shaded. In a preferred embodiment, a soluble form of BCR3 is provided wherein the signal peptide is deleted or preferably naturally cleaved, and the transmembrane domain is deleted, inactivated, or BCR3 is truncated to exclude the transmembrane domain.

Figure 16 shows the amino acid sequence of BCR3p1 and BCR3p2.

Figure 17 shows the relative amount of expression of BCR3 in various samples of breast cancer tissue (dark bars) and many normal tissue types (light bars).

Figure 18 shows an embodiment of a nucleic acid (mRNA) which includes a sequence which encodes a breast cancer protein provided herein, BCQ8. The start and stop codon are underlined.

Figure 19 shows an embodiment of an open reading frame of a nucleic acid encoding BCQ8, wherein the start and stop codons are underlined.

Figure 20 shows an embodiment of an amino acid sequence of BCQ8. The signal peptide is underlined twice and the transmembrane domain is underlined. In a preferred embodiment, a soluble form of BCQ8 is provided wherein the signal peptide is deleted, and the transmembrane domain is deleted, inactivated, or BCQ8 is truncated on either end as desired, to exclude the transmembrane domain.

Figure 21 shows the amino acid sequence of BCQ8p1 and BCQ8p2.

Figure 22 shows the relative amount of expression of BCQ8 in various samples of breast cancer tissue (dark bars) and many normal tissue types (light bars).

Figure 23 shows an embodiment of a nucleic acid (mRNA) which includes a sequence which encodes a differentially expressed protein provided herein, human BCQ5.

Figure 24 shows an embodiment of an open reading frame of a nucleic acid encoding human BCQ5.

Figure 25 shows embodiments of amino acid sequences of BCQ5 by providing an alignment wherein human is above mouse which is above rat.

Figure 26 shows the amino acid sequence of BCQ5p1, BCQ5p2 and BCQ5p3.

Figure 27 shows the relative amount of expression of BCQ5 in various samples of breast cancer tissue, colorectal cancer tissue, angiogenesis models wherein tubes are formed from endothelial cells, and normal tissue types.

Figure 28 shows an embodiment of a nucleic acid (mRNA) which includes a sequence which encodes a differentially expressed protein provided herein, mouse BCQ5.

Figure 29 shows an embodiment of an open reading frame of a nucleic acid encoding mouse BCQ5.

Figure 30 shows an embodiment of a nucleic acid (partial mRNA) which includes a sequence which encodes a differentially expressed protein provided herein, rat BCQ5.

Figure 31 shows an embodiment of a partial open reading frame of a nucleic acid encoding rat BCQ5.

Figure 32 shows an embodiment of a nucleic acid (mRNA) which includes a sequence which encodes a breast cancer protein provided herein, BCH1. Start and stop codons are underlined.

Figure 33 shows an embodiment of an open reading frame of a nucleic acid encoding BCH1, wherein start and stop codons are underlined.

Figure 34 shows an embodiment of an amino acid sequence of BCH1. In a preferred embodiment, isolated BCH1 excludes the signal sequence, amino acids 1-19.

Figure 35 shows the amino acid sequence of BCH1p1 and BCH1p2 (below).

Figure 36 shows the relative amount of expression of BCH1 in various samples of breast cancer tissue (dark bars) and many normal tissue types (light bars).

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Figure 37 shows a graph correlating expression of estrogen receptor (ER) (vertical bar) and BCH1 (horizontal bar) with localization to the nucleus (diamonds) or the cytoplasm (squares).

Figures 38A and 38B show breast carcinoma tissue specimens having high levels of BCH1 expression as indicated by anti-BCH1 antibodies.

Figures 39A and 39B show samples taken from the tissues shown in Figures 38A and 38B respectively, wherein anti-estrogen receptor antibodies indicate that estrogen receptor is localized exclusively to the cytoplasm.

Figures 40A and 40B show breast carcinoma tissue specimens having low levels of BCH1 expression as indicated by anti-BCH1 antibodies.

Figures 41A and 41B show samples taken from the tissues shown in Figures 40A and 40B respectively, wherein anti-estrogen receptor antibodies indicate that estrogen receptor in low BCH1 expression tissue does not correlate with estrogen receptor localization.

Figure 42 shows an embodiment of a nucleic acid (mRNA) which includes a sequence which encodes a breast cancer protein provided herein, BCN1. Start and stop codons are shaded, and Accession number AA419622 sequence is underlined.

Figure 43 shows an embodiment of an amino acid sequence of BCN1. A putative transmembrane domain is predicted to be at least at approximately positions 201-217 and 67-83. The protein may be a type IIIa membrane protein and may have additional transmembrane domains.

Figure 44 shows the amino acid sequence of BCN1p1 and BCN1p2.

Figure 45 shows the relative amount of expression of BCN1 in various samples of breast cancer tissue (dark bars), colon cancer tissue (light bars) and many normal tissue types (medium light dotted bars).

Figures 46A-46D show an embodiment of a nucleic acid (mRNA) which includes a sequence which encodes a breast cancer protein provided herein, BCN2. Start and stop codons are shaded, and Accession number AA428090 sequence is underlined.

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Figure 47 shows an embodiment of an amino acid sequence of BCN2.

Figure 48 shows the relative amount of expression of BCN2 in various samples of breast cancer tissue (dark bars), colon cancer tissue (light bars) and many normal tissue types (medium light dotted bars).

Figures 49A-49B show an embodiment of a nucleic acid (mRNA) which includes a sequence which encodes a breast cancer protein provided herein, BCN5. Start and stop codons are shaded, and Accession number R51309 sequence is underlined.

Figure 50 shows an embodiment of an amino acid sequence of BCN5. A putative signal sequence is shaded and preferred sequence is underlined.

Figure 51 shows the relative amount of expression of BCN5 in various samples of breast cancer tissue (dark bars), colon cancer tissue (light bars) and many normal tissue types (medium light dotted bars).

Figure 52 shows an embodiment of a nucleic acid (mRNA) which includes a sequence which encodes a breast cancer protein provided herein, BCO2. Start and stop codons are underlined.

Figure 53 shows an embodiment of an open reading frame of a nucleic acid encoding BCO2.

Figure 54 shows an embodiment of an amino acid sequence of BCO2.

Figure 55 shows an alignment of amino acids for human BCO2 above mouse BCO2.

Figure 56 shows the relative amount of expression of BCO2 in various samples of breast cancer tissue (dark bars) and many normal tissue types (light bars).

Figure 57 shows an embodiment of a nucleic acid (mRNA) which includes a sequence which encodes a breast cancer protein provided herein, BCX2. Start and stop codons are underlined.

Figure 58 shows an embodiment of an open reading frame of a nucleic acid encoding BCX2.

Figure 59 shows an embodiment of an amino acid sequence of BCX2.

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Figure 60 shows the relative amount of expression of BCX2 in various samples of breast cancer tissue (dark bars) and many normal tissue types (light bars).

Figure 61 shows the relative amount of expression of BCX2 in various samples of colon cancer tissue (light bars), non-cancer cell lines (dark bars) and many cell lines (medium light bars).

Figure 62 shows an embodiment of a nucleic acid (mRNA) which includes a sequence which encodes a breast cancer protein provided herein, BCX3. Start and stop codons are underlined.

Figure 63 shows an embodiment of an open reading frame of a nucleic acid encoding BCX3.

Figure 64 shows an embodiment of an amino acid sequence of BCX3.

Figure 65 shows the relative amount of expression of BCX3 in various samples of breast cancer tissue (dark bars) and many normal tissue types (light bars).

Figure 66 shows an embodiment of a nucleic acid (mRNA) which includes a sequence which encodes a breast cancer protein provided herein, BCA2. Start and stop codons are shaded, and Accession number D12485 sequence is underlined.

Figure 67 shows an embodiment of an amino acid sequence of BCA2.

Figure 68 shows an embodiment of a nucleic acid (mRNA) which includes a sequence which encodes a breast cancer protein provided herein, BCR2. Start and stop codons are shaded, and Accession number AA609773 sequence is shown underlined.

Figure 69 shows an embodiment of an amino acid sequence of BCR2.

Figure 70 shows the relative amount of expression of BCR2 in various samples of breast cancer tissue.

Figure 71 shows the relative amount of expression of BCR2 in various samples of normal tissue types (light bars).

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Figure 72 shows an embodiment of a nucleic acid which includes a sequence which encodes a breast cancer protein provided herein, BCJ7.

Figure 73 shows an embodiment of a nucleic acid which includes a sequence which encodes a breast cancer protein provided herein, BCY3.

DETAILED DESCRIPTION OF THE INVENTION

The present invention provides novel methods for diagnosis and prognosis evaluation for breast cancer, as well as methods for screening for compositions which modulate breast cancer. In one aspect, the expression levels of genes are determined in different patient samples for which either diagnosis or prognosis information is desired, to provide expression profiles. An expression profile of a particular sample is essentially a "fingerprint" of the state of the sample; while two states may have any particular gene similarly expressed, the evaluation of a number of genes simultaneously allows the generation of a gene expression profile that is unique to the state of the cell. That is, normal tissue may be distinguished from breast cancer tissue, and within breast cancer tissue, different prognosis states (good or poor long term survival prospects, for example) may be determined. By comparing expression profiles of breast tissue in known different states, information regarding which genes are important (including both up- and down-regulation of genes) in each of these states is obtained. The identification of sequences that are differentially expressed in breast cancer versus normal breast tissue, as well as differential expression resulting in different prognostic outcomes, allows the use of this information in a number of ways. For example, the evaluation of a particular treatment regime may be evaluated: does a chemotherapeutic drug act to improve the long-term prognosis in a particular patient. Similarly, diagnosis may be done or confirmed by comparing patient samples with the known expression profiles. Furthermore, these gene expression profiles (or individual genes) allow screening of drug candidates with an eye to mimicking or altering a particular expression profile; for example, screening can be done for drugs that suppress the breast cancer expression profile or convert a poor prognosis profile to a better prognosis profile. This may be done by making biochips comprising sets of the important breast cancer genes, which can then be used in these screens. These methods can also be done on the protein basis; that is, protein expression levels of the breast cancer proteins can be evaluated for diagnostic and prognostic purposes or to screen candidate agents. In addition, the breast cancer nucleic acid sequences can be administered for gene therapy purposes, including the administration of antisense nucleic acids, or the breast cancer proteins (including antibodies and other modulators thereof) administered as therapeutic drugs.

Thus the present invention provides nucleic acid and protein sequences that are differentially expressed in breast cancer, breast cancer, herein termed "breast cancer sequences". As outlined below, breast cancer sequences include those that are up-regulated (i.e. expressed at a higher level) in breast cancer, as well as those that are down-regulated (i.e. expressed at a lower level) in breast cancer. In a preferred embodiment, the breast cancer sequences are from humans; however, as will be appreciated by those in the art, breast cancer sequences from other organisms may be useful in animal models of disease and drug evaluation; thus, other breast cancer sequences are provided, from vertebrates, including mammals, including rodents (rats, mice, hamsters, guinea pigs, etc.), primates, farm animals (including sheep, goats, pigs, cows, horses, etc). Breast cancer sequences from other organisms may be obtained using the techniques outlined below.

Breast cancer sequences can include both nucleic acid and amino acid sequences. In a preferred embodiment, the breast cancer sequences are recombinant nucleic acids. By the term "recombinant nucleic acid" herein is meant nucleic acid, originally formed in vitro, in general, by the manipulation of nucleic acid by polymerases and endonucleases, in a form not normally found in nature. Thus an isolated nucleic acid, in a linear form, or an expression vector formed in vitro by ligating DNA molecules that are not normally joined, are both considered recombinant for the purposes of this invention. It is understood that once a recombinant nucleic acid is made and reintroduced into a host cell or organism, it will replicate non-recombinantly, i.e. using the in vivo cellular machinery of the host cell rather than in vitro manipulations; however, such nucleic acids, once produced recombinantly, although subsequently replicated non-recombinantly, are still considered recombinant for the purposes of the invention.

Similarly, a "recombinant protein" is a protein made using recombinant techniques, i.e. through the expression of a recombinant nucleic acid as depicted above. A recombinant protein is distinguished from naturally occurring protein by at least one or more characteristics. For example, the protein may be isolated or purified away from some or all of the proteins and compounds with which it is normally associated in its wild type host, and thus may be substantially pure. For example, an isolated protein is unaccompanied by at least some of the material with which it is normally associated in its natural state, preferably constituting at least about 0.5%, more preferably at least about 5% by weight of the total protein in a given sample. A substantially pure protein comprises at least about 75% by weight of the total protein, with at least about 80% being preferred, and at least about 90% being particularly preferred. The definition includes the production of a breast cancer protein from one organism in a

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different organism or host cell. Alternatively, the protein may be made at a significantly higher concentration than is normally seen, through the use of an inducible promoter or high expression promoter, such that the protein is made at increased concentration levels. Alternatively, the protein may be in a form not normally found in nature, as in the addition of an epitope tag or amino acid substitutions, insertions and deletions, as discussed below.

In a preferred embodiment, the breast cancer sequences are nucleic acids. As will be appreciated by those in the art and is more fully outlined below, breast cancer sequences are useful in a variety of applications, including diagnostic applications, which will detect naturally occurring nucleic acids, as well as screening applications; for example, biochips comprising nucleic acid probes to the breast cancer sequences can be generated. In the broadest sense, then, by "nucleic acid" or "oligonucleotide" or grammatical equivalents herein means at least two nucleotides covalently linked together. A nucleic acid of the present invention will generally contain phosphodiester bonds, although in some cases, as outlined below, nucleic acid analogs are included that may have alternate backbones, comprising, for example, phosphoramidate (Beaucage et al., Tetrahedron 49(10):1925 (1993) and references therein; Letsinger, J. Org. Chem. 35:3800 (1970); Sprinzl et al., Eur. J. Biochem. 81:579 (1977); Letsinger et al., Nucl. Acids Res. 14:3487 (1986); Sawai et al, Chem. Lett. 805 (1984), Letsinger et al., J. Am. Chem. Soc. 110:4470 (1988); and Pauwels et al., Chemica Scripta 26:141 91986)), phosphorothioate (Mag et al., Nucleic Acids Res. 19:1437 (1991); and U.S. Patent No. 5,644,048), phosphorodithioate (Briu et al., J. Am. Chem. Soc. 111:2321 (1989), Omethylphophoroamidite linkages (see Eckstein, Oligonucleotides and Analogues: A Practical Approach, Oxford University Press), and peptide nucleic acid backbones and linkages (see Egholm, J. Am. Chem. Soc. 114:1895 (1992); Meier et al., Chem. Int. Ed. Engl. 31:1008 (1992); Nielsen, Nature, 365:566 (1993); Carlsson et al., Nature 380:207 (1996), all of which are incorporated by reference). Other analog nucleic acids include those with positive backbones (Denpcy et al., Proc. Natl. Acad. Sci. USA 92:6097 (1995); non-ionic backbones (U.S. Patent Nos. 5,386,023, 5,637,684, 5,602,240, 5,216,141 and 4,469,863; Kiedrowshi et al., Angew. Chem. Intl. Ed. English 30:423 (1991); Letsinger et al., J. Am. Chem. Soc. 110:4470 (1988); Letsinger et al., Nucleoside & Nucleotide 13:1597 (1994); Chapters 2 and 3, ASC Symposium Series 580, "Carbohydrate Modifications in Antisense Research", Ed. Y.S. Sanghui and P. Dan Cook; Mesmaeker et al., Bioorganic & Medicinal Chem. Lett. 4:395 (1994); Jeffs et al., J. Biomolecular NMR 34:17 (1994); Tetrahedron Lett. 37:743 (1996)) and nonribose backbones, including those described in U.S. Patent Nos. 5,235,033 and 5,034,506, and Chapters 6 and 7, ASC Symposium Series 580, "Carbohydrate Modifications in Antisense Research", Ed. Y.S. Sanghui and P. Dan Cook. Nucleic acids containing one or more carbocyclic sugars are also

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included within one definition of nucleic acids (see Jenkins et al., Chem. Soc. Rev. (1995) pp169-176). Several nucleic acid analogs are described in Rawls, C & E News June 2, 1997 page 35. All of these references are hereby expressly incorporated by reference. These modifications of the ribose-phosphate backbone may be done for a variety of reasons, for example to increase the stability and half-life of such molecules in physiological environments or as probes on a biochip.

As will be appreciated by those in the art, all of these nucleic acid analogs may find use in the present invention. In addition, mixtures of naturally occurring nucleic acids and analogs can be made; alternatively, mixtures of different nucleic acid analogs, and mixtures of naturally occurring nucleic acids and analogs may be made.

Particularly preferred are peptide nucleic acids (PNA) which includes peptide nucleic acid analogs. These backbones are substantially non-ionic under neutral conditions, in contrast to the highly charged phosphodiester backbone of naturally occurring nucleic acids. This results in two advantages. First, the PNA backbone exhibits improved hybridization kinetics. PNAs have larger changes in the melting temperature (Tm) for mismatched versus perfectly matched basepairs. DNA and RNA typically exhibit a 2-4°C drop in Tm for an internal mismatch. With the non-ionic PNA backbone, the drop is closer to 7-9°C. Similarly, due to their non-ionic nature, hybridization of the bases attached to these backbones is relatively insensitive to salt concentration. In addition, PNAs are not degraded by cellular enzymes, and thus can be more stable.

The nucleic acids may be single stranded or double stranded, as specified, or contain portions of both double stranded or single stranded sequence. As will be appreciated by those in the art, the depiction of a single strand ("Watson") also defines the sequence of the other strand ("Crick"); thus the sequences described herein also includes the complement of the sequence. The nucleic acid may be DNA, both genomic and cDNA, RNA or a hybrid, where the nucleic acid contains any combination of deoxyribo- and ribo-nucleotides, and any combination of bases, including uracil, adenine, thymine, cytosine, guanine, inosine, xanthine hypoxanthine, isocytosine, isoguanine, etc. As used herein, the term "nucleoside" includes nucleotides and nucleoside and nucleotide analogs, and modified nucleosides such as amino modified nucleosides. In addition, "nucleoside" includes non-naturally occurring analog structures. Thus for example the individual units of a peptide nucleic acid, each containing a base, are referred to herein as a nucleoside.

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A breast cancer sequence can be initially identified by substantial nucleic acid and/or amino acid sequence homology to the breast cancer sequences outlined herein. Such homology can be based upon the overall nucleic acid or amino acid sequence, and is generally determined as outlined below, using either homology programs or hybridization conditions.

The breast cancer sequences of the invention can be identified as follows. Samples of normal and tumor tissue are applied to biochips comprising nucleic acid probes. The samples are first microdissected, if applicable, and treated as is know in the art for the preparation of mRNA. Suitable biochips are commercially available, for example from Affymetrix. Gene expression profiles as described herein are generated, and the data analyzed.

In a preferred embodiment, the genes showing changes in expression as between normal and disease states are compared to genes expressed in other normal tissues, including, but not limited to lung, heart, brain, liver, breast, kidney, muscle, prostate, small intestine, large intestine, spleen, bone, and placenta. In a preferred embodiment, those genes identified during the breast cancer screen that are expressed in any significant amount in other tissues are removed from the profile, although in some embodiments, this is not necessary. That is, when screening for drugs, it is preferable that the target be disease specific, to minimize possible side effects.

In a preferred embodiment, breast cancer sequences are those that are up-regulated in breast cancer; that is, the expression of these genes is higher in breast carcinoma as compared to normal breast tissue. "Up-regulation" as used herein means at least about a two-fold change, preferably at least about a three fold change, with at least about five-fold or higher being preferred. All accession numbers herein are for the GenBank sequence database and the sequences of the accession numbers are hereby expressly incorporated by reference. GenBank is known in the art, see, e.g., Benson, DA, et al., Nucleic Acids Research 26:1-7 (1998) and http://www.ncbi.nlm.nih.gov/. In addition, these genes were found to be expressed in a limited relative amount or not at all in heart, brain, lung, liver, kidney, prostate, small intestine, testes, fibroblasts and spleen.

In a preferred embodiment, breast cancer sequences are those that are down-regulated in breast cancer; that is, the expression of these genes is lower in breast carcinoma as compared to normal breast tissue. "Down-regulation" as used herein means at least about a two-fold change, preferably at least about a three fold change, with at least about five-fold or higher being preferred.

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Breast cancer proteins of the present invention may be classified as secreted proteins, transmembrane proteins or intracellular proteins. In a preferred embodiment the breast cancer protein is an intracellular protein. Intracellular proteins are involved in all aspects of cellular function and replication (including, for example, signaling pathways); aberrant expression of such proteins results in unregulated or disregulated cellular processes. For example, many intracellular proteins have enzymatic activity such as protein kinase activity, protein phosphatase activity, protease activity, nucleotide cyclase activity, polymerase activity and the like. Intracellular proteins also serve as docking proteins that are involved in organizing complexes of proteins, or targeting proteins to various subcellular localizations, and are involved in maintaining the structural integrity of organelles.

An increasingly appreciated concept in characterizing intracellular proteins is the presence in the proteins of one or more motifs for which defined functions have been attributed. In addition to the highly conserved sequences found in the enzymatic domain of proteins, highly conserved sequences have been identified in proteins that are involved in protein-protein interaction. For example, Srchomology-2 (SH2) domains bind tyrosine-phosphorylated targets in a sequence dependent manner. PTB domains, which are distinct from SH2 domains, also bind tyrosine phosphorylated targets. SH3 domains bind to proline-rich targets. In addition, PH domains, tetratricopeptide repeats and WD domains to name only a few, have been shown to mediate protein-protein interactions. Some of these may also be involved in binding to phospholipids or other second messengers. As will be appreciated by one of ordinary skill in the art, these motifs can be identified on the basis of primary sequence; thus, an analysis of the sequence of proteins may provide insight into both the enzymatic potential of the molecule and/or molecules with which the protein may associate.

In a preferred embodiment, the breast cancer sequences are transmembrane proteins.

Transmembrane proteins are molecules that span the phospholipid bilayer of a cell. They may have an intracellular domain, an extracellular domain, or both. The intracellular domains of such proteins may have a number of functions including those already described for intracellular proteins. For example, the intracellular domain may have enzymatic activity and/or may serve as a binding site for additional proteins. Frequently the intracellular domain of transmembrane proteins serves both roles. For example certain receptor tyrosine kinases have both protein kinase activity and SH2 domains. In addition, autophosphorylation of tyrosines on the receptor molecule itself, creates binding sites for additional SH2 domain containing proteins.

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Transmembrane proteins may contain from one to many transmembrane domains. For example, receptor tyrosine kinases, certain cytokine receptors, receptor guanylyl cyclases and receptor serine/threonine protein kinases contain a single transmembrane domain. However, various other proteins including channels and adenylyl cyclases contain numerous transmembrane domains. Many important cell surface receptors are classified as "seven transmembrane domain" proteins, as they contain 7 membrane spanning regions. Important transmembrane protein receptors include, but are not limited to insulin receptor, insulin-like growth factor receptor, human growth hormone receptor, glucose transporters, transferrin receptor, epidermal growth factor receptor, low density lipoprotein receptor, epidermal growth factor receptor, interleukin receptors, e.g. IL-1 receptor, IL-2 receptor, etc.

Characteristics of transmembrane domains include approximately 20 consecutive hydrophobic amino acids that may be followed by charged amino acids. Therefore, upon analysis of the amino acid sequence of a particular protein, the localization and number of transmembrane domains within the protein may be predicted.

The extracellular domains of transmembrane proteins are diverse; however, conserved motifs are found repeatedly among various extracellular domains. Conserved structure and/or functions have been ascribed to different extracellular motifs. For example, cytokine receptors are characterized by a cluster of cysteines and a WSXWS (W= tryptophan, S= serine, X=any amino acid) motif. Immunoglobulin-like domains are highly conserved. Mucin-like domains may be involved in cell adhesion and leucine-rich repeats participate in protein-protein interactions. It is understood that the transmembrane domains may be removed to create soluble proteins herein.

Many extracellular domains are involved in binding to other molecules. In one aspect, extracellular domains are receptors. Factors that bind the receptor domain include circulating ligands, which may be peptides, proteins, or small molecules such as adenosine and the like. For example, growth factors such as EGF, FGF and PDGF are circulating growth factors that bind to their cognate receptors to initiate a variety of cellular responses. Other factors include cytokines, mitogenic factors, neurotrophic factors and the like. Extracellular domains also bind to cell-associated molecules. In this respect, they mediate cell-cell interactions. Cell-associated ligands can be tethered to the cell for example via a glycosylphosphatidylinositol (GPI) anchor, or may themselves be transmembrane proteins. Extracellular domains also associate with the extracellular matrix and contribute to the maintenance of the cell structure.

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Breast cancer proteins that are transmembrane are particularly preferred in the present invention as they are good targets for immunotherapeutics, as are described herein. In addition, as outlined below, transmembrane proteins can be also useful in imaging modalities.

In a preferred embodiment, the breast cancer proteins are secreted proteins; the secretion of which can be either constitutive or regulated. These proteins have a signal peptide or signal sequence that targets the molecule to the secretory pathway. Secreted proteins are involved in numerous physiological events; by virtue of their circulating nature, they serve to transmit signals to various other cell types. The secreted protein may function in an autocrine manner (acting on the cell that secreted the factor), a paracrine manner (acting on cells in close proximity to the cell that secreted the factor) or an endocrine manner (acting on cells at a distance). Thus secreted molecules find use in modulating or altering numerous aspects of physiology. Breast cancer proteins that are secreted proteins are particularly preferred in the present invention as they serve as good targets for diagnostic markers, for example for blood tests.

A breast cancer sequence is initially identified by substantial nucleic acid and/or amino acid sequence homology to the breast cancer sequences outlined herein. Such homology can be based upon the overall nucleic acid or amino acid sequence, and is generally determined as outlined below, using either homology programs or hybridization conditions.

As used herein, a nucleic acid is a "breast cancer nucleic acid" if the overall homology of the nucleic acid sequence to the nucleic acid sequences encoding the amino acid sequences of the figures is preferably greater than about 75%, more preferably greater than about 80%, even more preferably greater than about 85% and most preferably greater than 90%. In some embodiments the homology will be as high as about 93 to 95 or 98%. Homology in this context means sequence similarity or identity, with identity being preferred. A preferred comparison for homology purposes is to compare the sequence containing sequencing errors to the correct sequence. This homology will be determined using standard techniques known in the art, including, but not limited to, the local homology algorithm of Smith & Waterman, Adv. Appl. Math. 2:482 (1981), by the homology alignment algorithm of Needleman & Wunsch, J. Mol. Biool. 48:443 (1970), by the search for similarity method of Pearson & Lipman, PNAS USA 85:2444 (1988), by computerized implementations of these algorithms (GAP, BESTFIT, FASTA, and TFASTA in the Wisconsin Genetics Software Package, Genetics Computer Group, 575 Science Drive, Madison, WI), the Best Fit sequence program described by

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Devereux et al., Nucl. Acid Res. 12:387-395 (1984), preferably using the default settings, or by inspection.

In a preferred embodiment, the sequences which are used to determine sequence identity or similarity are selected from the sequences set forth in the figures. In one embodiment the sequences utilized herein are those set forth in the figures. In another embodiment, the sequences are naturally occurring allelic variants of the sequences set forth in the figures. In another embodiment, the sequences are sequence variants as further described herein.

One example of a useful algorithm is PILEUP. PILEUP creates a multiple sequence alignment from a group of related sequences using progressive, pairwise alignments. It can also plot a tree showing the clustering relationships used to create the alignment. PILEUP uses a simplification of the progressive alignment method of Feng & Doolittle, J. Mol. Evol. 35:351-360 (1987); the method is similar to that described by Higgins & Sharp CABIOS 5:151-153 (1989). Useful PILEUP parameters including a default gap weight of 3.00, a default gap length weight of 0.10, and weighted end gaps.

Another example of a useful algorithm is the BLAST algorithm, described in Altschul et al., J. Mol. Biol. 215, 403-410, (1990) and Karlin et al., PNAS USA 90:5873-5787 (1993). A particularly useful BLAST program is the WU-BLAST-2 program which was obtained from Altschul et al., Methods in Enzymology, 266: 460-480 (1996); http://blast.wustl/edu/blast/ READE.html]. WU-BLAST-2 uses several search parameters, most of which are set to the default values. The adjustable parameters are set with the following values: overlap span =1, overlap fraction = 0.125, word threshold (T) = 11. The HSP S and HSP S2 parameters are dynamic values and are established by the program itself depending upon the composition of the particular sequence and composition of the particular database against which the sequence of interest is being searched; however, the values may be adjusted to increase sensitivity. A % amino acid sequence identity value is determined by the number of matching identical residues divided by the total number of residues of the "longer" sequence in the aligned region. The "longer" sequence is the one having the most actual residues in the aligned region (gaps introduced by WU-Blast-2 to maximize the alignment score are ignored).

Thus, "percent (%) nucleic acid sequence identity" is defined as the percentage of nucleotide residues in a candidate sequence that are identical with the nucleotide residues of the se. A preferred method utilizes the BLASTN module of WU-BLAST-2 set to the default parameters, with overlap span and overlap fraction set to 1 and 0.125, respectively.

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The alignment may include the introduction of gaps in the sequences to be aligned. In addition, for sequences which contain either more or fewer nucleosides than those of the figures, it is understood that the percentage of homology will be determined based on the number of homologous nucleosides in relation to the total number of nucleosides. Thus, for example, homology of sequences shorter than those of the sequences identified herein and as discussed below, will be determined using the number of nucleosides in the shorter sequence.

In one embodiment, the nucleic acid homology is determined through hybridization studies. Thus, for example, nucleic acids which hybridize under high stringency to the nucleic acid sequences which encode the proteins identified in the figures, or their complements, are considered a breast cancer sequence. High stringency conditions are known in the art; see for example Maniatis et al., Molecular Cloning: A Laboratory Manual, 2d Edition, 1989, and Short Protocols in Molecular Biology, ed. Ausubel, et al., both of which are hereby incorporated by reference. Stringent conditions are sequence-dependent and will be different in different circumstances. Longer sequences hybridize specifically at higher temperatures. An extensive guide to the hybridization of nucleic acids is found in Tijssen, Techniques in Biochemistry and Molecular Biology--Hybridization with Nucleic Acid Probes, "Overview of principles of hybridization and the strategy of nucleic acid assays" (1993). Generally, stringent conditions are selected to be about 5-10°C lower than the thermal melting point (Tm) for the specific sequence at a defined ionic strength pH. The Tm is the temperature (under defined ionic strength, pH and nucleic acid concentration) at which 50% of the probes complementary to the target hybridize to the target sequence at equilibrium (as the target sequences are present in excess, at Tm, 50% of the probes are occupied at equilibrium). Stringent conditions will be those in which the salt concentration is less than about 1.0 M sodium ion, typically about 0.01 to 1.0 M sodium ion concentration (or other salts) at pH 7.0 to 8.3 and the temperature is at least about 30°C for short probes (e.g. 10 to 50 nucleotides) and at least about 60°C for long probes (e.g. greater than 50 nucleotides). Stringent conditions may also be achieved with the addition of destabilizing agents such as formamide.

In another embodiment, less stringent hybridization conditions are used; for example, moderate or low stringency conditions may be used, as are known in the art; see Maniatis and Ausubel, supra, and Tijssen, supra.

In addition, the breast cancer nucleic acid sequences of the invention are fragments of larger genes, i.e. they are nucleic acid segments. "Genes" in this context includes coding regions, non-coding

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regions, and mixtures of coding and non-coding regions. Accordingly, as will be appreciated by those in the art, using the sequences provided herein, additional sequences of the breast cancer genes can be obtained, using techniques well known in the art for cloning either longer sequences or the full length sequences; see Maniatis et al., and Ausubel, et al., supra, hereby expressly incorporated by reference.

Once the breast cancer nucleic acid is identified, it can be cloned and, if necessary, its constituent parts recombined to form the entire breast cancer nucleic acid. Once isolated from its natural source, e.g., contained within a plasmid or other vector or excised therefrom as a linear nucleic acid segment, the recombinant breast cancer nucleic acid can be further-used as a probe to identify and isolate other breast cancer nucleic acids, for example additional coding regions. It can also be used as a "precursor" nucleic acid to make modified or variant breast cancer nucleic acids and proteins.

The breast cancer nucleic acids of the present invention are used in several ways. In a first embodiment, nucleic acid probes to the breast cancer nucleic acids are made and attached to biochips to be used in screening and diagnostic methods, as outlined below, or for administration, for example for gene therapy and/or antisense applications. Alternatively, the breast cancer nucleic acids that include coding regions of breast cancer proteins can be put into expression vectors for the expression of breast cancer proteins, again either for screening purposes or for administration to a patient.

In a preferred embodiment, nucleic acid probes to breast cancer nucleic acids (both the nucleic acid sequences encoding peptides outlined in the figures and/or the complements thereof) are made. The nucleic acid probes attached to the biochip are designed to be substantially complementary to the breast cancer nucleic acids, i.e. the target sequence (either the target sequence of the sample or to other probe sequences, for example in sandwich assays), such that hybridization of the target sequence and the probes of the present invention occurs. As outlined below, this complementarity need not be perfect; there may be any number of base pair mismatches which will interfere with hybridization between the target sequence and the single stranded nucleic acids of the present invention. However, if the number of mutations is so great that no hybridization can occur under even the least stringent of hybridization conditions, the sequence is not a complementary target sequence. Thus, by "substantially complementary" herein is meant that the probes are sufficiently complementary to the target sequences to hybridize under normal reaction conditions, particularly high stringency conditions, as outlined herein.

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A nucleic acid probe is generally single stranded but can be partially single and partially double stranded. The strandedness of the probe is dictated by the structure, composition, and properties of the target sequence. In general, the nucleic acid probes range from about 8 to about 100 bases long, with from about 10 to about 80 bases being preferred, and from about 30 to about 50 bases being particularly preferred. That is, generally whole genes are not used. In some embodiments, much longer nucleic acids can be used, up to hundreds of bases.

In a preferred embodiment, more than one probe per sequence is used, with either overlapping probes or probes to different sections of the target being used. That is, two, three, four or more probes, with three being preferred, are used to build in a redundancy for a particular target. The probes can be overlapping (i.e. have some sequence in common), or separate.

As will be appreciated by those in the art, nucleic acids can be attached or immobilized to a solid support in a wide variety of ways. By "immobilized" and grammatical equivalents herein is meant the association or binding between the nucleic acid probe and the solid support is sufficient to be stable under the conditions of binding, washing, analysis, and removal as outlined below. The binding can be covalent or non-covalent. By "non-covalent binding" and grammatical equivalents herein is meant one or more of either electrostatic, hydrophilic, and hydrophobic interactions. Included in non-covalent binding is the covalent attachment of a molecule, such as, streptavidin to the support and the non-covalent binding of the biotinylated probe to the streptavidin. By "covalent binding" and grammatical equivalents herein is meant that the two moieties, the solid support and the probe, are attached by at least one bond, including sigma bonds, pi bonds and coordination bonds. Covalent bonds can be formed directly between the probe and the solid support or can be formed by a cross linker or by inclusion of a specific reactive group on either the solid support or the probe or both molecules. Immobilization may also involve a combination of covalent and non-covalent interactions.

In general, the probes are attached to the biochip in a wide variety of ways, as will be appreciated by those in the art. As described herein, the nucleic acids can either be synthesized first, with subsequent attachment to the biochip, or can be directly synthesized on the biochip.

The biochip comprises a suitable solid substrate. By "substrate" or "solid support" or other grammatical equivalents herein is meant any material that can be modified to contain discrete individual sites appropriate for the attachment or association of the nucleic acid probes and is amenable to at least one detection method. As will be appreciated by those in the art, the number of

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possible substrates are very large, and include, but are not limited to, glass and modified or functionalized glass, plastics (including acrylics, polystyrene and copolymers of styrene and other materials, polypropylene, polyethylene, polybutylene, polyurethanes, TeflonJ, etc.), polysaccharides, nylon or nitrocellulose, resins, silica or silica-based materials including silicon and modified silicon, carbon, metals, inorganic glasses, plastics, etc. In general, the substrates allow optical detection and do not appreciably fluorescese. A preferred substrate is described in copending application entitled A Reusable Low Fluorescent Plastic Biochip, U.S. Application Serial No. 09/270,214, filed March 15, 1999, herein incorporated by reference in its entirety.

Generally the substrate is planar, although as will be appreciated by those in the art, other configurations of substrates may be used as well. For example, the probes may be placed on the inside surface of a tube, for flow-through sample analysis to minimize sample volume. Similarly, the substrate may be flexible, such as a flexible foam, including closed cell foams made of particular plastics.

In a preferred embodiment, the surface of the biochip and the probe may be derivatized with chemical functional groups for subsequent attachment of the two. Thus, for example, the biochip is derivatized with a chemical functional group including, but not limited to, amino groups, carboxy groups, oxo groups and thiol groups, with amino groups being particularly preferred. Using these functional groups, the probes can be attached using functional groups on the probes. For example, nucleic acids containing amino groups can be attached to surfaces comprising amino groups, for example using linkers as are known in the art; for example, homo-or hetero-bifunctional linkers as are well known (see 1994 Pierce Chemical Company catalog, technical section on cross-linkers, pages 155-200, incorporated herein by reference). In addition, in some cases, additional linkers, such as alkyl groups (including substituted and heteroalkyl groups) may be used.

In this embodiment, the oligonucleotides are synthesized as is known in the art, and then attached to the surface of the solid support. As will be appreciated by those skilled in the art, either the 5' or 3' terminus may be attached to the solid support, or attachment may be via an internal nucleoside.

In an additional embodiment, the immobilization to the solid support may be very strong, yet non-covalent. For example, biotinylated oligonucleotides can be made, which bind to surfaces covalently coated with streptavidin, resulting in attachment.

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Alternatively, the oligonucleotides may be synthesized on the surface, as is known in the art. For example, photoactivation techniques utilizing photopolymerization compounds and techniques are used. In a preferred embodiment, the nucleic acids can be synthesized in situ, using well known photolithographic techniques, such as those described in WO 95/25116; WO 95/35505; U.S. Patent Nos. 5,700,637 and 5,445,934; and references cited within, all of which are expressly incorporated by reference; these methods of attachment form the basis of the Affimetrix GeneChip™ technology.

In a preferred embodiment, breast cancer nucleic acids encoding breast cancer proteins are used to make a variety of expression vectors to express breast cancer proteins which can then be used in screening assays, as described below. The expression vectors may be either self-replicating extrachromosomal vectors or vectors which integrate into a host genome. Generally, these expression vectors include transcriptional and translational regulatory nucleic acid operably linked to the nucleic acid encoding the breast cancer protein. The term "control sequences" refers to DNA sequences necessary for the expression of an operably linked coding sequence in a particular host organism. The control sequences that are suitable for prokaryotes, for example, include a promoter, optionally an operator sequence, and a ribosome binding site. Eukaryotic cells are known to utilize promoters, polyadenylation signals, and enhancers.

Nucleic acid is "operably linked" when it is placed into a functional relationship with another nucleic acid sequence. For example, DNA for a presequence or secretory leader is operably linked to DNA for a polypeptide if it is expressed as a preprotein that participates in the secretion of the polypeptide; a promoter or enhancer is operably linked to a coding sequence if it affects the transcription of the sequence; or a ribosome binding site is operably linked to a coding sequence if it is positioned so as to facilitate translation. Generally, "operably linked" means that the DNA sequences being linked are contiguous, and, in the case of a secretory leader, contiguous and in reading phase. However, enhancers do not have to be contiguous. Linking is accomplished by ligation at convenient restriction sites. If such sites do not exist, the synthetic oligonucleotide adaptors or linkers are used in accordance with conventional practice. The transcriptional and translational regulatory nucleic acid will generally be appropriate to the host cell used to express the breast cancer protein; for example, transcriptional and translational regulatory nucleic acid sequences from *Bacillus* are preferably used to express the breast cancer protein in *Bacillus*. Numerous types of appropriate expression vectors, and suitable regulatory sequences are known in the art for a variety of host cells.

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In general, the transcriptional and translational regulatory sequences may include, but are not limited to, promoter sequences, ribosomal binding sites, transcriptional start and stop sequences, translational start and stop sequences, and enhancer or activator sequences. In a preferred embodiment, the regulatory sequences include a promoter and transcriptional start and stop sequences.

Promoter sequences encode either constitutive or inducible promoters. The promoters may be either naturally occurring promoters or hybrid promoters. Hybrid promoters, which combine elements of more than one promoter, are also known in the art, and are useful in the present invention.

In addition, the expression vector may comprise additional elements. For example, the expression vector may have two replication systems, thus allowing it to be maintained in two organisms, for example in mammalian or insect cells for expression and in a procaryotic host for cloning and amplification. Furthermore, for integrating expression vectors, the expression vector contains at least one sequence homologous to the host cell genome, and preferably two homologous sequences which flank the expression construct. The integrating vector may be directed to a specific locus in the host cell by selecting the appropriate homologous sequence for inclusion in the vector. Constructs for integrating vectors are well known in the art.

In addition, in a preferred embodiment, the expression vector contains a selectable marker gene to allow the selection of transformed host cells. Selection genes are well known in the art and will vary with the host cell used.

The breast cancer proteins of the present invention are produced by culturing a host cell transformed with an expression vector containing nucleic acid encoding a breast cancer protein, under the appropriate conditions to induce or cause expression of the breast cancer protein. The conditions appropriate for breast cancer protein expression will vary with the choice of the expression vector and the host cell, and will be easily ascertained by one skilled in the art through routine experimentation. For example, the use of constitutive promoters in the expression vector will require optimizing the growth and proliferation of the host cell, while the use of an inducible promoter requires the appropriate growth conditions for induction. In addition, in some embodiments, the timing of the harvest is important. For example, the baculoviral systems used in insect cell expression are lytic viruses, and thus harvest time selection can be crucial for product yield.

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Appropriate host cells include yeast, bacteria, archaebacteria, fungi, and insect and animal cells, including mammalian cells. Of particular interest are *Drosophila melangaster* cells, *Saccharomyces cerevisiae* and other yeasts, *E. coli*, *Bacillus subtilis*, Sf9 cells, C129 cells, 293 cells, *Neurospora*, BHK, CHO, COS, HeLa cells, THP1 cell line (a macrophage cell line) and human cells and cell lines.

In a preferred embodiment, the breast cancer proteins are expressed in mammalian cells. Mammalian expression systems are also known in the art, and include retroviral systems. A preferred expression vector system is a retroviral vector system such as is generally described in PCT/US97/01019 and PCT/US97/01048, both of which are hereby expressly incorporated by reference. Of particular use as mammalian promoters are the promoters from mammalian viral genes, since the viral genes are often highly expressed and have a broad host range. Examples include the SV40 early promoter, mouse mammary tumor virus LTR promoter, adenovirus major late promoter, herpes simplex virus promoter, and the CMV promoter. Typically, transcription termination and polyadenylation sequences recognized by mammalian cells are regulatory regions located 3' to the translation stop codon and thus, together with the promoter elements, flank the coding sequence. Examples of transcription terminator and polyadenylytion signals include those derived form SV40.

The methods of introducing exogenous nucleic acid into mammalian hosts, as well as other hosts, is well known in the art, and will vary with the host cell used. Techniques include dextran-mediated transfection, calcium phosphate precipitation, polybrene mediated transfection, protoplast fusion, electroporation, viral infection, encapsulation of the polynucleotide(s) in liposomes, and direct microinjection of the DNA into nuclei.

In a preferred embodiment, breast cancer proteins are expressed in bacterial systems. Bacterial expression systems are well known in the art. Promoters from bacteriophage may also be used and are known in the art. In addition, synthetic promoters and hybrid promoters are also useful; for example, the tac promoter is a hybrid of the trp and lac promoter sequences. Furthermore, a bacterial promoter can include naturally occurring promoters of non-bacterial origin that have the ability to bind bacterial RNA polymerase and initiate transcription. In addition to a functioning promoter sequence, an efficient ribosome binding site is desirable. The expression vector may also include a signal peptide sequence that provides for secretion of the breast cancer protein in bacteria. The protein is either secreted into the growth media (gram-positive bacteria) or into the periplasmic space, located between the inner and outer membrane of the cell (gram-negative bacteria). The bacterial expression vector may also include a selectable marker gene to allow for the selection of bacterial strains that

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have been transformed. Suitable selection genes include genes which render the bacteria resistant to drugs such as ampicillin, chloramphenicol, erythromycin, kanamycin, neomycin and tetracycline. Selectable markers also include biosynthetic genes, such as those in the histidine, tryptophan and leucine biosynthetic pathways. These components are assembled into expression vectors. Expression vectors for bacteria are well known in the art, and include vectors for *Bacillus subtilis*, *E. coli*, *Streptococcus cremoris*, and *Streptococcus lividans*, among others. The bacterial expression vectors are transformed into bacterial host cells using techniques well known in the art, such as calcium chloride treatment, electroporation, and others.

In one embodiment, breast cancer proteins are produced in insect cells. Expression vectors for the transformation of insect cells, and in particular, baculovirus-based expression vectors, are well known in the art.

In a preferred embodiment, breast cancer protein is produced in yeast cells. Yeast expression systems are well known in the art, and include expression vectors for *Saccharomyces cerevisiae*, Candida albicans and C. maltosa, Hansenula polymorpha, Kluyveromyces fragilis and K. lactis, Pichia guillerimondii and P. pastoris, Schizosaccharomyces pombe, and Yarrowia lipolytica.

The breast cancer protein may also be made as a fusion protein, using techniques well known in the art. Thus, for example, for the creation of monoclonal antibodies, if the desired epitope is small, the breast cancer protein may be fused to a carrier protein to form an immunogen. Alternatively, the breast cancer protein may be made as a fusion protein to increase expression, or for other reasons. For example, when the breast cancer protein is a breast cancer peptide, the nucleic acid encoding the peptide may be linked to other nucleic acid for expression purposes.

In one embodiment, the breast cancer nucleic acids, proteins and antibodies of the invention are labeled. By "labeled" herein is meant that a compound has at least one element, isotope or chemical compound attached to enable the detection of the compound. In general, labels fall into three classes: a) isotopic labels, which may be radioactive or heavy isotopes; b) immune labels, which may be antibodies or antigens; and c) colored or fluorescent dyes. The labels may be incorporated into the breast cancer nucleic acids, proteins and antibodies at any position. For example, the label should be capable of producing, either directly or indirectly, a detectable signal. The detectable moiety may be a radioisotope, such as ³H, ¹⁴C, ³²P, ³⁵S, or ¹²⁵I, a fluorescent or chemiluminescent compound, such as fluorescein isothiocyanate, rhodamine, or luciferin, or an enzyme, such as alkaline phosphatase, beta-

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galactosidase or horseradish peroxidase. Any method known in the art for conjugating the antibody to the label may be employed, including those methods described by Hunter et al., <u>Nature</u>, <u>144</u>:945 (1962); David et al., <u>Biochemistry</u>, <u>13</u>:1014 (1974); Pain et al., <u>J. Immunol. Meth.</u>, <u>40</u>:219 (1981); and Nygren, <u>J. Histochem. and Cytochem.</u>, <u>30</u>:407 (1982).

Accordingly, the present invention also provides breast cancer protein sequences. A breast cancer protein of the present invention may be identified in several ways. "Protein" in this sense includes proteins, polypeptides, and peptides. As will be appreciated by those in the art, the nucleic acid sequences of the invention can be used to generate protein sequences. There are a variety of ways to do this, including cloning the entire gene and verifying its frame and amino acid sequence, or by comparing it to known sequences to search for homology to provide a frame, assuming the breast cancer protein has homology to some protein in the database being used. Generally, the nucleic acid sequences are input into a program that will search all three frames for homology. This is done in a preferred embodiment using the following NCBI Advanced BLAST parameters. The program is blastx or blastn. The database is nr. The input data is as "Sequence in FASTA format". The organism list is "none". The "expect" is 10; the filter is default. The "descriptions" is 500, the "alignments" is 500, and the "alignment view" is pairwise. The "Query Genetic Codes" is standard (1). The matrix is BLOSUM62; gap existence cost is 11, per residue gap cost is 1; and the lambda ratio is .85 default. This results in the generation of a putative protein sequence.

Also included within one embodiment of breast cancer proteins are amino acid variants of the naturally occurring sequences, as determined herein. Preferably, the variants are preferably greater than about 75% homologous to the wild-type sequence, more preferably greater than about 80%, even more preferably greater than about 85% and most preferably greater than 90%. In some embodiments the homology will be as high as about 93 to 95 or 98%. As for nucleic acids, homology in this context means sequence similarity or identity, with identity being preferred. This homology will be determined using standard techniques known in the art as are outlined above for the nucleic acid homologies.

Breast cancer proteins of the present invention may be shorter or longer than the wild type amino acid sequences. Thus, in a preferred embodiment, included within the definition of breast cancer proteins are portions or fragments of the wild type sequences. herein. In addition, as outlined above, the breast cancer nucleic acids of the invention may be used to obtain additional coding regions, and thus additional protein sequence, using techniques known in the art.

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In a preferred embodiment, the breast cancer proteins are derivative or variant breast cancer proteins as compared to the wild-type sequence. That is, as outlined more fully below, the derivative breast cancer peptide will contain at least one amino acid substitution, deletion or insertion, with amino acid substitutions being particularly preferred. The amino acid substitution, insertion or deletion may occur at any residue within the breast cancer peptide.

Also included in an embodiment of breast cancer proteins of the present invention are amino acid sequence variants. These variants fall into one or more of three classes: substitutional, insertional or deletional variants. These variants ordinarily are prepared by site specific mutagenesis of nucleotides in the DNA encoding the breast cancer protein, using cassette or PCR mutagenesis or other techniques well known in the art, to produce DNA encoding the variant, and thereafter expressing the DNA in recombinant cell culture as outlined above. However, variant breast cancer protein fragments having up to about 100-150 residues may be prepared by in vitro synthesis using established techniques. Amino acid sequence variants are characterized by the predetermined nature of the variation, a feature that sets them apart from naturally occurring allelic or interspecies variation of the breast cancer protein amino acid sequence. The variants typically exhibit the same qualitative biological activity as the naturally occurring analogue, although variants can also be selected which have modified characteristics as will be more fully outlined below.

While the site or region for introducing an amino acid sequence variation is predetermined, the mutation per se need not be predetermined. For example, in order to optimize the performance of a mutation at a given site, random mutagenesis may be conducted at the target codon or region and the expressed breast cancer variants screened for the optimal combination of desired activity. Techniques for making substitution mutations at predetermined sites in DNA having a known sequence are well known, for example, M13 primer mutagenesis and PCR mutagenesis. Screening of the mutants is done using assays of breast cancer protein activities.

Amino acid substitutions are typically of single residues; insertions usually will be on the order of from about 1 to 20 amino acids, although considerably larger insertions may be tolerated. Deletions range from about 1 to about 20 residues, although in some cases deletions may be much larger.

Substitutions, deletions, insertions or any combination thereof may be used to arrive at a final derivative. Generally these changes are done on a few amino acids to minimize the alteration of the molecule. However, larger changes may be tolerated in certain circumstances. When small

alterations in the characteristics of the breast cancer protein are desired, substitutions are generally made in accordance with the following chart:

Chart

		Chart I
	Original Residue	Exemplary Substitutions
5	Ala	Ser
	Arg	Lys
	Asn	Gln, His
	Asp	Glu
	Cys	Ser
10	Gln	Asn
	Glu	Asp
	Gly	Pro
	His	Asn, Gln
	lle	Leu, Val
15	Leu	lle, Val
4	Lys	Arg, Gln, Glu
J	Met	Leu, Ile
5	Phe	Met, Leu, Tyr
The state of the s	Ser	Thr
20	Thr	Ser
ij	Trp	Tyr
in Es	Tyr	Trp, Phe
É	Val	Ile, Leu
interior in the second		

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Substantial changes in function or immunological identity are made by selecting substitutions that are less conservative than those shown in Chart I. For example, substitutions may be made which more significantly affect: the structure of the polypeptide backbone in the area of the alteration, for example the alpha-helical or beta-sheet structure; the charge or hydrophobicity of the molecule at the target site; or the bulk of the side chain. The substitutions which in general are expected to produce the greatest changes in the polypeptide's properties are those in which (a) a hydrophilic residue, e.g. seryl or threonyl is substituted for (or by) a hydrophobic residue, e.g. leucyl, isoleucyl, phenylalanyl, valyl or alanyl; (b) a cysteine or proline is substituted for (or by) any other residue; (c) a residue having an electropositive side chain, e.g. lysyl, arginyl, or histidyl, is substituted for (or by) an electronegative residue, e.g. glutamyl or aspartyl; or (d) a residue having a bulky side chain, e.g. phenylalanine, is substituted for (or by) one not having a side chain, e.g. glycine.

The variants typically exhibit the same qualitative biological activity and will elicit the same immune response as the naturally-occurring analogue, although variants also are selected to modify the characteristics of the breast cancer proteins as needed. Alternatively, the variant may be designed

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such that the biological activity of the breast cancer protein is altered. For example, glycosylation sites may be altered or removed.

Covalent modifications of breast cancer polypeptides are included within the scope of this invention. One type of covalent modification includes reacting targeted amino acid residues of a breast cancer polypeptide with an organic derivatizing agent that is capable of reacting with selected side chains or the N-or C-terminal residues of a breast cancer polypeptide. Derivatization with bifunctional agents is useful, for instance, for crosslinking breast cancer to a water-insoluble support matrix or surface for use in the method for purifying anti-breast cancer antibodies or screening assays, as is more fully described below. Commonly used crosslinking agents include, e.g., 1,1-bis(diazoacetyl)-2-phenylethane, glutaraldehyde, N-hydroxysuccinimide esters, for example, esters with 4-azidosalicylic acid, homobifunctional imidoesters, including disuccinimidyl esters such as 3,3'-dithiobis(succinimidyl-propionate), bifunctional maleimides such as bis-N-maleimido-1,8-octane and agents such as methyl-3-[(p-azidophenyl)dithio]propioimidate.

Other modifications include deamidation of glutaminyl and asparaginyl residues to the corresponding glutamyl and aspartyl residues, respectively, hydroxylation of proline and lysine, phosphorylation of hydroxyl groups of seryl, threonyl or tyrosyl residues, methylation of the α -amino groups of lysine, arginine, and histidine side chains [T.E. Creighton, Proteins: Structure and Molecular Properties, W.H. Freeman & Co., San Francisco, pp. 79-86 (1983)], acetylation of the N-terminal amine, and amidation of any C-terminal carboxyl group.

Another type of covalent modification of the breast cancer polypeptide included within the scope of this invention comprises altering the native glycosylation pattern of the polypeptide. "Altering the native glycosylation pattern" is intended for purposes herein to mean deleting one or more carbohydrate moieties found in native sequence breast cancer polypeptide, and/or adding one or more glycosylation sites that are not present in the native sequence breast cancer polypeptide.

Addition of glycosylation sites to breast cancer polypeptides may be accomplished by altering the amino acid sequence thereof. The alteration may be made, for example, by the addition of, or substitution by, one or more serine or threonine residues to the native sequence breast cancer polypeptide (for O-linked glycosylation sites). The breast cancer amino acid sequence may optionally be altered through changes at the DNA level, particularly by mutating the DNA encoding the breast

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cancer polypeptide at preselected bases such that codons are generated that will translate into the desired amino acids.

Another means of increasing the number of carbohydrate moieties on the breast cancer polypeptide is by chemical or enzymatic coupling of glycosides to the polypeptide. Such methods are described in the art, e.g., in WO 87/05330 published 11 September 1987, and in Aplin and Wriston, breast cancer Crit. Rev. Biochem., pp. 259-306 (1981).

Removal of carbohydrate moieties present on the breast cancer polypeptide may be accomplished chemically or enzymatically or by mutational substitution of codons encoding for amino acid residues that serve as targets for glycosylation. Chemical deglycosylation techniques are known in the art and described, for instance, by Hakimuddin, et al., Arch. Biochem. Biophys., 259:52 (1987) and by Edge et al., Anal. Biochem., 118:131 (1981). Enzymatic cleavage of carbohydrate moieties on polypeptides can be achieved by the use of a variety of endo-and exo-glycosidases as described by Thotakura et al., Meth. Enzymol., 138:350 (1987).

Another type of covalent modification of breast cancer comprises linking the breast cancer polypeptide to one of a variety of nonproteinaceous polymers, e.g., polyethylene glycol, polypropylene glycol, or polyoxyalkylenes, in the manner set forth in U.S. Patent Nos. 4,640,835; 4,496,689; 4,301,144; 4,670,417; 4,791,192 or 4,179,337.

Breast cancer polypeptides of the present invention may also be modified in a way to form chimeric molecules comprising a breast cancer polypeptide fused to another, heterologous polypeptide or amino acid sequence. In one embodiment, such a chimeric molecule comprises a fusion of a breast cancer polypeptide with a tag polypeptide which provides an epitope to which an anti-tag antibody can selectively bind. The epitope tag is generally placed at the amino-or carboxyl-terminus of the breast cancer polypeptide. The presence of such epitope-tagged forms of a breast cancer polypeptide can be detected using an antibody against the tag polypeptide. Also, provision of the epitope tag enables the breast cancer polypeptide to be readily purified by affinity purification using an anti-tag antibody or another type of affinity matrix that binds to the epitope tag. In an alternative embodiment, the chimeric molecule may comprise a fusion of a breast cancer polypeptide with an immunoglobulin or a particular region of an immunoglobulin. For a bivalent form of the chimeric molecule, such a fusion could be to the Fc region of an IgG molecule.

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Various tag polypeptides and their respective antibodies are well known in the art. Examples include poly-histidine (poly-his) or poly-histidine-glycine (poly-his-gly) tags; the flu HA tag polypeptide and its antibody 12CA5 [Field et al., Mol. Cell. Biol., 8:2159-2165 (1988)]; the c-myc tag and the 8F9, 3C7, 6E10, G4, B7 and 9E10 antibodies thereto [Evan et al., Molecular and Cellular Biology, 5:3610-3616 (1985)]; and the Herpes Simplex virus glycoprotein D (gD) tag and its antibody [Paborsky et al., Protein Engineering, 3(6):547-553 (1990)]. Other tag polypeptides include the Flag-peptide [Hopp et al., BioTechnology, 6:1204-1210 (1988)]; the KT3 epitope peptide [Martin et al., Science, 255:192-194 (1992)]; tubulin epitope peptide [Skinner et al., J. Biol. Chem., 266:15163-15166 (1991)]; and the T7 gene 10 protein peptide tag [Lutz-Freyermuth et al., Proc. Natl. Acad. Sci. USA, 87:6393-6397 (1990)].

Also included with the definition of breast cancer protein in one embodiment are other breast cancer proteins of the breast cancer family, and breast cancer proteins from other organisms, which are cloned and expressed as outlined below. Thus, probe or degenerate polymerase chain reaction (PCR) primer sequences may be used to find other related breast cancer proteins from humans or other organisms. As will be appreciated by those in the art, particularly useful probe and/or PCR primer sequences include the unique areas of the breast cancer nucleic acid sequence. As is generally known in the art, preferred PCR primers are from about 15 to about 35 nucleotides in length, with from about 20 to about 30 being preferred, and may contain inosine as needed. The conditions for the PCR reaction are well known in the art.

In addition, as is outlined herein, breast cancer proteins can be made that are longer than those depicted in the figures, for example, by the elucidation of additional sequences, the addition of epitope or purification tags, the addition of other fusion sequences, etc.

Breast cancer proteins may also be identified as being encoded by breast cancer nucleic acids. Thus, breast cancer proteins are encoded by nucleic acids that will hybridize to the sequences of the sequence listings, or their complements, as outlined herein.

In a preferred embodiment, when the breast cancer protein is to be used to generate antibodies, for example for immunotherapy, the breast cancer protein should share at least one epitope or determinant with the full length protein. By "epitope" or "determinant" herein is meant a portion of a protein which will generate and/or bind an antibody or T-cell receptor in the context of MHC. Thus, in most instances, antibodies made to a smaller breast cancer protein will be able to bind to the full length protein. In a preferred embodiment, the epitope is unique; that is, antibodies generated to a

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unique epitope show little or no cross-reactivity. In a preferred embodiment, the epitope is selected from BCH1p1, BCH1p2, BCR3p1, BCR3p2, BCQ8p1, BCQ8p2, BCQ5p1, BCQ5p2, BCN1p1, BCN1p2, wherein BCH1p1 and BCH1p2 are preferred. In one embodiment the epitope or fragment of BCH1p1 is conjugated to KLH or BSA.

In one embodiment, the term "antibody" includes antibody fragments, as are known in the art, including Fab, Fab₂, single chain antibodies (Fv for example), chimeric antibodies, etc., either produced by the modification of whole antibodies or those synthesized de novo using recombinant DNA technologies.

Methods of preparing polyclonal antibodies are known to the skilled artisan. Polyclonal antibodies can be raised in a mammal, for example, by one or more injections of an immunizing agent and, if desired, an adjuvant. Typically, the immunizing agent and/or adjuvant will be injected in the mammal by multiple subcutaneous or intraperitoneal injections. The immunizing agent may include the BCH1 or fragment thereof or a fusion protein thereof. It may be useful to conjugate the immunizing agent to a protein known to be immunogenic in the mammal being immunized. Examples of such immunogenic proteins include but are not limited to keyhole limpet hemocyanin, serum albumin, bovine thyroglobulin, and soybean trypsin inhibitor. Examples of adjuvants which may be employed include Freund's complete adjuvant and MPL-TDM adjuvant (monophosphoryl Lipid A, synthetic trehalose dicorynomycolate). The immunization protocol may be selected by one skilled in the art without undue experimentation.

The antibodies may, alternatively, be monoclonal antibodies. Monoclonal antibodies may be prepared using hybridoma methods, such as those described by Kohler and Milstein, Nature, 256:495 (1975). In a hybridoma method, a mouse, hamster, or other appropriate host animal, is typically immunized with an immunizing agent to elicit lymphocytes that produce or are capable of producing antibodies that will specifically bind to the immunizing agent. Alternatively, the lymphocytes may be immunized in vitro. The immunizing agent will typically include the BCH1 polypeptide or fragment thereof or a fusion protein thereof. Generally, either peripheral blood lymphocytes ("PBLs") are used if cells of human origin are desired, or spleen cells or lymph node cells are used if non-human mammalian sources are desired. The lymphocytes are then fused with an immortalized cell line using a suitable fusing agent, such as polyethylene glycol, to form a hybridoma cell [Goding, Monoclonal Antibodies: Principles and Practice, Academic Press, (1986) pp. 59-103]. Immortalized cell lines are usually transformed mammalian cells, particularly myeloma cells of rodent, bovine and human origin. Usually, rat or

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mouse myeloma cell lines are employed. The hybridoma cells may be cultured in a suitable culture medium that preferably contains one or more substances that inhibit the growth or survival of the unfused, immortalized cells. For example, if the parental cells lack the enzyme hypoxanthine guanine phosphoribosyl transferase (HGPRT or HPRT), the culture medium for the hybridomas typically will include hypoxanthine, aminopterin, and thymidine ("HAT medium"), which substances prevent the growth of HGPRT-deficient cells.

In one embodiment, the antibodies are bispecific antibodies. Bispecific antibodies are monoclonal, preferably human or humanized, antibodies that have binding specificities for at least two different antigens. In the present case, one of the binding specificities is for the breast cancer protein or a fragment thereof, the other one is for any other antigen, and preferably for a cell-surface protein or receptor or receptor subunit, preferably one that is tumor specific.

In a preferred embodiment, the antibodies to breast cancer are capable of reducing or eliminating the biological function of breast cancer, as is described below. That is, the addition of anti-breast cancer antibodies (either polyclonal or preferably monoclonal) to breast cancer (or cells containing breast cancer) may reduce or eliminate the breast cancer activity. Generally, at least a 25% decrease in activity is preferred, with at least about 50% being particularly preferred and about a 95-100% decrease being especially preferred.

In a preferred embodiment the antibodies to the breast cancer proteins are humanized antibodies. Humanized forms of non-human (e.g., murine) antibodies are chimeric molecules of immunoglobulins, immunoglobulin chains or fragments thereof (such as Fv, Fab, Fab', F(ab')₂ or other antigen-binding subsequences of antibodies) which contain minimal sequence derived from non-human immunoglobulin. Humanized antibodies include human immunoglobulins (recipient antibody) in which residues form a complementary determining region (CDR) of the recipient are replaced by residues from a CDR of a non-human species (donor antibody) such as mouse, rat or rabbit having the desired specificity, affinity and capacity. In some instances, Fv framework residues of the human immunoglobulin are replaced by corresponding non-human residues. Humanized antibodies may also comprise residues which are found neither in the recipient antibody nor in the imported CDR or framework sequences. In general, the humanized antibody will comprise substantially all of at least one, and typically two, variable domains, in which all or substantially all of the CDR regions correspond to those of a non-human immunoglobulin and all or substantially all of the FR regions are those of a human immunoglobulin consensus sequence. The humanized antibody optimally also will

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comprise at least a portion of an immunoglobulin constant region (Fc), typically that of a human immunoglobulin [Jones et al., <u>Nature</u>, <u>321</u>:522-525 (1986); Riechmann et al., <u>Nature</u>, <u>332</u>:323-329 (1988); and Presta, Curr. Op. Struct. Biol., <u>2</u>:593-596 (1992)].

Methods for humanizing non-human antibodies are well known in the art. Generally, a humanized antibody has one or more amino acid residues introduced into it from a source which is non-human. These non-human amino acid residues are often referred to as import residues, which are typically taken from an import variable domain. Humanization can be essentially performed following the method of Winter and co-workers [Jones et al., Nature, 321:522-525 (1986); Riechmann et al., Nature, 332:323-327 (1988); Verhoeyen et al., Science, 239:1534-1536 (1988)], by substituting rodent CDRs or CDR sequences for the corresponding sequences of a human antibody. Accordingly, such humanized antibodies are chimeric antibodies (U.S. Patent No. 4,816,567), wherein substantially less than an intact human variable domain has been substituted by the corresponding sequence from a non-human species. In practice, humanized antibodies are typically human antibodies in which some CDR residues and possibly some FR residues are substituted by residues from analogous sites in rodent antibodies.

Human antibodies can also be produced using various techniques known in the art, including phage display libraries [Hoogenboom and Winter, J. Mol. Biol., 227:381 (1991); Marks et al., J. Mol. Biol., 222:581 (1991)]. The techniques of Cole et al. and Boerner et al. are also available for the preparation of human monoclonal antibodies (Cole et al., Monoclonal Antibodies and Cancer Therapy, Alan R. Liss, p. 77 (1985) and Boerner et al., J. Immunol., 147(1):86-95 (1991)]. Similarly, human antibodies can be made by introducing of human immunoglobulin loci into transgenic animals, e.g., mice in which the endogenous immunoglobulin genes have been partially or completely inactivated. Upon challenge, human antibody production is observed, which closely resembles that seen in humans in all respects, including gene rearrangement, assembly, and antibody repertoire. This approach is described, for example, in U.S. Patent Nos. 5,545,807; 5,545,806; 5,569,825; 5,625,126; 5,633,425; 5,661,016, and in the following scientific publications: Marks et al., Bio/Technology 10, 779-783 (1992); Lonberg et al., Nature 368, 856-859 (1994); Morrison, Nature 368, 812-13 (1994); Fishwild et al., Nature Biotechnology 14, 845-51 (1996); Neuberger, Nature Biotechnology 14, 826 (1996); Lonberg and Huszar, Intern. Rev. Immunol. 13 65-93 (1995).

By immunotherapy is meant treatment of breast cancer with an antibody raised against breast cancer proteins. As used herein, immunotherapy can be passive or active. Passive immunotherapy as

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defined herein is the passive transfer of antibody to a recipient (patient). Active immunization is the induction of antibody and/or T-cell responses in a recipient (patient). Induction of an immune response is the result of providing the recipient with an antigen to which antibodies are raised. As appreciated by one of ordinary skill in the art, the antigen may be provided by injecting a polypeptide against which antibodies are desired to be raised into a recipient, or contacting the recipient with a nucleic acid capable of expressing the antigen and under conditions for expression of the antigen.

In a preferred embodiment the breast cancer proteins against which antibodies are raised are secreted proteins as described above. Without being bound by theory, antibodies used for treatment, bind and prevent the secreted protein from binding to its receptor, thereby inactivating the secreted breast cancer protein.

In another preferred embodiment, the breast cancer protein to which antibodies are raised is a transmembrane protein. Without being bound by theory, antibodies used for treatment, bind the extracellular domain of the breast cancer protein and prevent it from binding to other proteins, such as circulating ligands or cell-associated molecules. The antibody may cause down-regulation of the transmembrane breast cancer protein. As will be appreciated by one of ordinary skill in the art, the antibody may be a competitive, non-competitive or uncompetitive inhibitor of protein binding to the extracellular domain of the breast cancer protein. The antibody is also an antagonist of the breast cancer protein. Further, the antibody prevents activation of the transmembrane breast cancer protein. In one aspect, when the antibody prevents the binding of other molecules to the breast cancer protein, the antibody prevents growth of the cell. The antibody also sensitizes the cell to cytotoxic agents, including, but not limited to TNF- α , TNF- β , IL-1, INF- γ and IL-2, or chemotherapeutic agents including 5FU, vinblastine, actinomycin D, cisplatin, methotrexate, and the like. In some instances the antibody belongs to a sub-type that activates serum complement when complexed with the transmembrane protein thereby mediating cytotoxicity. Thus, breast cancer is treated by administering to a patient antibodies directed against the transmembrane breast cancer protein.

In another preferred embodiment, the antibody is conjugated to a therapeutic moiety. In one aspect the therapeutic moiety is a small molecule that modulates the activity of the breast cancer protein. In another aspect the therapeutic moiety modulates the activity of molecules associated with or in close proximity to the breast cancer protein. The therapeutic moiety may inhibit enzymatic activity such as protease or protein kinase activity associated with breast cancer.

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In a preferred embodiment, the therapeutic moiety may also be a cytotoxic agent. In this method, targeting the cytotoxic agent to tumor tissue or cells, results in a reduction in the number of afflicted cells, thereby reducing symptoms associated with breast cancer. Cytotoxic agents are numerous and varied and include, but are not limited to, cytotoxic drugs or toxins or active fragments of such toxins. Suitable toxins and their corresponding fragments include diptheria A chain, exotoxin A chain, ricin A chain, abrin A chain, curcin, crotin, phenomycin, enomycin and the like. Cytotoxic agents also include radiochemicals made by conjugating radioisotopes to antibodies raised against breast cancer proteins, or binding of a radionuclide to a chelating agent that has been covalently attached to the antibody. Targeting the therapeutic moiety to transmembrane breast cancer proteins not only serves to increase the local concentration of therapeutic moiety in the breast cancer afflicted area, but also serves to reduce deleterious side effects that may be associated with the therapeutic moiety.

The breast cancer antibodies of the invention specifically bind to breast cancer proteins. By "specifically bind" herein is meant that the antibodies bind to the protein with a binding constant in the range of at least 10⁻⁴- 10⁻⁶ M⁻¹, with a preferred range being 10⁻⁷ - 10⁻⁹ M⁻¹.

In a preferred embodiment, the breast cancer protein is purified or isolated after expression. Breast cancer proteins may be isolated or purified in a variety of ways known to those skilled in the art depending on what other components are present in the sample. Standard purification methods include electrophoretic, molecular, immunological and chromatographic techniques, including ion exchange, hydrophobic, affinity, and reverse-phase HPLC chromatography, and chromatofocusing. For example, the breast cancer protein may be purified using a standard anti-breast cancer antibody column. Ultrafiltration and diafiltration techniques, in conjunction with protein concentration, are also useful. For general guidance in suitable purification techniques, see Scopes, R., Protein Purification, Springer-Verlag, NY (1982). The degree of purification necessary will vary depending on the use of the breast cancer protein. In some instances no purification will be necessary.

Once expressed and purified if necessary, the breast cancer proteins and nucleic acids are useful in a number of applications.

In one aspect, the expression levels of genes are determined for different cellular states in the breast cancer phenotype; that is, the expression levels of genes in normal breast tissue and in breast cancer tissue (and in some cases, for varying severities of breast cancer that relate to prognosis, as outlined below) are evaluated to provide expression profiles. An expression profile of a particular cell state or

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point of development is essentially a "fingerprint" of the state; while two states may have any particular gene similarly expressed, the evaluation of a number of genes simultaneously allows the generation of a gene expression profile that is unique to the state of the cell. By comparing expression profiles of cells in different states, information regarding which genes are important (including both up- and down-regulation of genes) in each of these states is obtained. Then, diagnosis may be done or confirmed: does tissue from a particular patient have the gene expression profile of normal or breast cancer tissue.

"Differential expression," or grammatical equivalents as used herein, refers to both qualitative as well as quantitative differences in the genes' temporal and/or cellular expression patterns within and among the cells. Thus, a differentially expressed gene can qualitatively have its expression altered, including an activation or inactivation, in, for example, normal versus breast cancer tissue. That is, genes may be turned on or turned off in a particular state, relative to another state. As is apparent to the skilled artisan, any comparison of two or more states can be made. Such a qualitatively regulated gene will exhibit an expression pattern within a state or cell type which is detectable by standard techniques in one such state or cell type, but is not detectable in both. Alternatively, the determination is quantitative in that expression is increased or decreased; that is, the expression of the gene is either upregulated, resulting in an increased amount of transcript, or downregulated, resulting in a decreased amount of transcript. The degree to which expression differs need only be large enough to quantify via standard characterization techniques as outlined below, such as by use of Affymetrix GeneChip™ expression arrays, Lockhart, Nature Biotechnology, 14:1675-1680 (1996), hereby expressly incorporated by reference. Other techniques include, but are not limited to, quantitative reverse transcriptase PCR, Northern analysis and RNase protection. As outlined above, preferably the change in expression (i.e. upregulation or downregulation) is at least about 50%, more preferably at least about 100%, more preferably at least about 150%, more preferably, at least about 200%, with from 300 to at least 1000% being especially preferred.

As will be appreciated by those in the art, this may be done by evaluation at either the gene transcript, or the protein level; that is, the amount of gene expression may be monitored using nucleic acid probes to the DNA or RNA equivalent of the gene transcript, and the quantification of gene expression levels, or, alternatively, the final gene product itself (protein) can be monitored, for example through the use of antibodies to the breast cancer protein and standard immunoassays (ELISAs,e tc.) or other techniques, including mass spectroscopy assays, 2D gel electrophoresis assays, etc. Thus, the

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proteins corresponding to breast cancer genes, i.e. those identified as being important in a breast cancer phenotype, can be evaluated in a breast cancer diagnostic test.

In a preferred embodiment, gene expression monitoring is done and a number of genes, i.e. an expression profile, is monitored simultaneously, although multiple protein expression monitoring can be done as well. Similarly, these assays may be done on an individual basis as well.

In this embodiment, the breast cancer nucleic acid probes are attached to biochips as outlined herein for the detection and quantification of breast cancer sequences in a particular cell. The assays are further described below in the example.

In a preferred embodiment nucleic acids encoding the breast cancer protein are detected. Although DNA or RNA encoding the breast cancer protein may be detected, of particular interest are methods wherein the mRNA encoding a breast cancer protein is detected. The presence of mRNA in a sample is an indication that the breast cancer gene has been transcribed to form the mRNA, and suggests that the protein is expressed. Probes to detect the mRNA can be any nucleotide/deoxynucleotide probe that is complementary to and base pairs with the mRNA and includes but is not limited to oligonucleotides, cDNA or RNA. Probes also should contain a detectable label, as defined herein. In one method the mRNA is detected after immobilizing the nucleic acid to be examined on a solid support such as nylon membranes and hybridizing the probe with the sample. Following washing to remove the non-specifically bound probe, the label is detected. In another method detection of the mRNA is performed in situ. In this method permeabilized cells or tissue samples are contacted with a detectably labeled nucleic acid probe for sufficient time to allow the probe to hybridize with the target mRNA. Following washing to remove the non-specifically bound probe, the label is detected. For example a digoxygenin labeled riboprobe (RNA probe) that is complementary to the mRNA encoding a breast cancer protein is detected by binding the digoxygenin with an anti-digoxygenin secondary antibody and developed with nitro blue tetrazolium and 5-bromo-4-chloro-3-indoyl phosphate.

In a preferred embodiment, any of the three classes of proteins as described herein (secreted, transmembrane or intracellular proteins) are used in diagnostic assays. The breast cancer proteins, antibodies, nucleic acids, modified proteins and cells containing breast cancer sequences are used in diagnostic assays. This can be done on an individual gene or corresponding polypeptide level. In a preferred embodiment, the expression profiles are used, preferably in conjunction with high throughput

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screening techniques to allow monitoring for expression profile genes and/or corresponding polypeptides.

As described and defined herein, breast cancer proteins, including intracellular, transmembrane or secreted proteins, find use as markers of breast cancer. Detection of these proteins in putative breast cancer tissue or patients allows for a determination or diagnosis of breast cancer. Numerous methods known to those of ordinary skill in the art find use in detecting breast cancer. In one embodiment, antibodies are used to detect breast cancer proteins. A preferred method separates proteins from a sample or patient by electrophoresis on a gel (typically a denaturing and reducing protein gel, but may be any other type of gel including isoelectric focusing gels and the like). Following separation of proteins, the breast cancer protein is detected by immunoblotting with antibodies raised against the breast cancer protein. Methods of immunoblotting are well known to those of ordinary skill in the art.

In another preferred method, antibodies to the breast cancer protein find use in <u>in situ</u> imaging techniques. In this method cells are contacted with from one to many antibodies to the breast cancer protein(s). Following washing to remove non-specific antibody binding, the presence of the antibody or antibodies is detected. In one embodiment the antibody is detected by incubating with a secondary antibody that contains a detectable label. In another method the primary antibody to the breast cancer protein(s) contains a detectable label. In another preferred embodiment each one of multiple primary antibodies contains a distinct and detectable label. This method finds particular use in simultaneous screening for a plurality of breast cancer proteins. As will be appreciated by one of ordinary skill in the art, numerous other histological imaging techniques are useful in the invention.

In a preferred embodiment the label is detected in a fluorometer which has the ability to detect and distinguish emissions of different wavelengths. In addition, a fluorescence activated cell sorter (FACS) can be used in the method.

In another preferred embodiment, antibodies find use in diagnosing breast cancer from blood samples. As previously described, certain breast cancer proteins are secreted/circulating molecules. Blood samples, therefore, are useful as samples to be probed or tested for the presence of secreted breast cancer proteins. Antibodies can be used to detect the breast cancer by any of the previously described immunoassay techniques including ELISA, immunoblotting (Western blotting), immunoprecipitation, BIACORE technology and the like, as will be appreciated by one of ordinary skill in the art.

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In a preferred embodiment, <u>in situ</u> hybridization of labeled breast cancer nucleic acid probes to tissue arrays is done. For example, arrays of tissue samples, including breast cancer tissue and/or normal tissue, are made. <u>In situ</u> hybridization as is known in the art can then be done.

It is understood that when comparing the fingerprints between an individual and a standard, the skilled artisan can make a diagnosis as well as a prognosis. It is further understood that the genes which indicate the diagnosis may differ from those which indicate the prognosis.

In a preferred embodiment, the breast cancer proteins, antibodies, nucleic acids, modified proteins and cells containing breast cancer sequences are used in prognosis assays. As above, gene expression profiles can be generated that correlate to breast cancer severity, in terms of long term prognosis. Again, this may be done on either a protein or gene level, with the use of genes being preferred. As above, the breast cancer probes are attached to biochips for the detection and quantification of breast cancer sequences in a tissue or patient. The assays proceed as outlined for diagnosis.

In a preferred embodiment, any of the three classes of proteins as described herein are used in drug screening assays. The breast cancer proteins, antibodies, nucleic acids, modified proteins and cells containing breast cancer sequences are used in drug screening assays or by evaluating the effect of drug candidates on a "gene expression profile" or expression profile of polypeptides. In a preferred embodiment, the expression profiles are used, preferably in conjunction with high throughput screening techniques to allow monitoring for expression profile genes after treatment with a candidate agent, Zlokarnik, et al., Science 279, 84-8 (1998), Heid, 1996 #69.

In a preferred embodiment, the breast cancer proteins, antibodies, nucleic acids, modified proteins and cells containing the native or modified breast cancer proteins are used in screening assays. That is, the present invention provides novel methods for screening for compositions which modulate the breast cancer phenotype. As above, this can be done on an individual gene level or by evaluating the effect of drug candidates on a "gene expression profile". In a preferred embodiment, the expression profiles are used, preferably in conjunction with high throughput screening techniques to allow monitoring for expression profile genes after treatment with a candidate agent, see Zlokarnik, supra.

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Having identified the differentially expressed genes herein, a variety of assays may be executed. In a preferred embodiment, assays may be run on an individual gene or protein level. That is, having identified a particular gene as up regulated in breast cancer, candidate bioactive agents may be screened to modulate this gene's response; preferably to down regulate the gene, although in some circumstances to up regulate the gene. "Modulation" thus includes both an increase and a decrease in gene expression. The preferred amount of modulation will depend on the original change of the gene expression in normal versus tumor tissue, with changes of at least 10%, preferably 50%, more preferably 100-300%, and in some embodiments 300-1000% or greater. Thus, if a gene exhibits a 4 fold increase in tumor compared to normal tissue, a decrease of about four fold is desired; a 10 fold decrease in tumor compared to normal tissue gives a 10 fold increase in expression for a candidate agent is desired.

As will be appreciated by those in the art, this may be done by evaluation at either the gene or the protein level; that is, the amount of gene expression may be monitored using nucleic acid probes and the quantification of gene expression levels, or, alternatively, the gene product itself can be monitored, for example through the use of antibodies to the breast cancer protein and standard immunoassays.

In a preferred embodiment, gene expression monitoring is done and a number of genes, i.e. an expression profile, is monitored simultaneously, although multiple protein expression monitoring can be done as well.

In this embodiment, the breast cancer nucleic acid probes are attached to biochips as outlined herein for the detection and quantification of breast cancer sequences in a particular cell. The assays are further described below.

Generally, in a preferred embodiment, a candidate bioactive agent is added to the cells prior to analysis. Moreover, screens are provided to identify a candidate bioactive agent which modulates breast cancer, modulates breast cancer proteins, binds to a breast cancer protein, or interferes between the binding of a breast cancer protein and an antibody.

The term "candidate bioactive agent" or "drug candidate" or grammatical equivalents as used herein describes any molecule, e.g., protein, oligopeptide, small organic molecule, polysaccharide, polynucleotide, etc., to be tested for bioactive agents that are capable of directly or indirectly altering either the breast cancer phenotype or the expression of a breast cancer sequence, including both

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nucleic acid sequences and protein sequences. In preferred embodiments, the bioactive agents modulate the expression profiles, or expression profile nucleic acids or proteins provided herein. In a particularly preferred embodiment, the candidate agent suppresses a breast cancer phenotype, for example to a normal breast tissue fingerprint. Similarly, the candidate agent preferably suppresses a severe breast cancer phenotype. Generally a plurality of assay mixtures are run in parallel with different agent concentrations to obtain a differential response to the various concentrations. Typically, one of these concentrations serves as a negative control, i.e., at zero concentration or below the level of detection.

Candidate agents encompass numerous chemical classes, though typically they are organic molecules, preferably small organic compounds having a molecular weight of more than 100 and less than about 2,500 daltons. Preferred small molecules are less than 2000, or less than 1500 or less than 1500 or less than 500 D. Candidate agents comprise functional groups necessary for structural interaction with proteins, particularly hydrogen bonding, and typically include at least an amine, carbonyl, hydroxyl or carboxyl group, preferably at least two of the functional chemical groups. The candidate agents often comprise cyclical carbon or heterocyclic structures and/or aromatic or polyaromatic structures substituted with one or more of the above functional groups. Candidate agents are also found among biomolecules including peptides, saccharides, fatty acids, steroids, purines, pyrimidines, derivatives, structural analogs or combinations thereof. Particularly preferred are peptides.

Candidate agents are obtained from a wide variety of sources including libraries of synthetic or natural compounds. For example, numerous means are available for random and directed synthesis of a wide variety of organic compounds and biomolecules, including expression of randomized oligonucleotides. Alternatively, libraries of natural compounds in the form of bacterial, fungal, plant and animal extracts are available or readily produced. Additionally, natural or synthetically produced libraries and compounds are readily modified through conventional chemical, physical and biochemical means. Known pharmacological agents may be subjected to directed or random chemical modifications, such as acylation, alkylation, esterification, amidification to produce structural analogs.

In a preferred embodiment, the candidate bioactive agents are proteins. By "protein" herein is meant at least two covalently attached amino acids, which includes proteins, polypeptides, oligopeptides and peptides. The protein may be made up of naturally occurring amino acids and peptide bonds, or synthetic peptidomimetic structures. Thus "amino acid", or "peptide residue", as used herein means

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both naturally occurring and synthetic amino acids. For example, homo-phenylalanine, citrulline and noreleucine are considered amino acids for the purposes of the invention. "Amino acid" also includes imino acid residues such as proline and hydroxyproline. The side chains may be in either the (R) or the (S) configuration. In the preferred embodiment, the amino acids are in the (S) or L-configuration. If non-naturally occurring side chains are used, non-amino acid substituents may be used, for example to prevent or retard in vivo degradations.

In a preferred embodiment, the candidate bioactive agents are naturally occurring proteins or fragments of naturally occurring proteins. Thus, for example, cellular extracts containing proteins, or random or directed digests of proteinaceous cellular extracts, may be used. In this way libraries of procaryotic and eucaryotic proteins may be made for screening in the methods of the invention. Particularly preferred in this embodiment are libraries of bacterial, fungal, viral, and mammalian proteins, with the latter being preferred, and human proteins being especially preferred.

In a preferred embodiment, the candidate bioactive agents are peptides of from about 5 to about 30 amino acids, with from about 5 to about 20 amino acids being preferred, and from about 7 to about 15 being particularly preferred. The peptides may be digests of naturally occurring proteins as is outlined above, random peptides, or "biased" random peptides. By "randomized" or grammatical equivalents herein is meant that each nucleic acid and peptide consists of essentially random nucleotides and amino acids, respectively. Since generally these random peptides (or nucleic acids, discussed below) are chemically synthesized, they may incorporate any nucleotide or amino acid at any position. The synthetic process can be designed to generate randomized proteins or nucleic acids, to allow the formation of all or most of the possible combinations over the length of the sequence, thus forming a library of randomized candidate bioactive proteinaceous agents.

In one embodiment, the library is fully randomized, with no sequence preferences or constants at any position. In a preferred embodiment, the library is biased. That is, some positions within the sequence are either held constant, or are selected from a limited number of possibilities. For example, in a preferred embodiment, the nucleotides or amino acid residues are randomized within a defined class, for example, of hydrophobic amino acids, hydrophilic residues, sterically biased (either small or large) residues, towards the creation of nucleic acid binding domains, the creation of cysteines, for cross-linking, prolines for SH-3 domains, serines, threonines, tyrosines or histidines for phosphorylation sites, etc., or to purines, etc.

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In a preferred embodiment, the candidate bioactive agents are nucleic acids, as defined above.

As described above generally for proteins, nucleic acid candidate bioactive agents may be naturally occurring nucleic acids, random nucleic acids, or "biased" random nucleic acids. For example, digests of procaryotic or eucaryotic genomes may be used as is outlined above for proteins.

In a preferred embodiment, the candidate bioactive agents are organic chemical moieties, a wide variety of which are available in the literature.

After the candidate agent has been added and the cells allowed to incubate for some period of time, the sample containing the target sequences to be analyzed is added to the biochip. If required, the target sequence is prepared using known techniques. For example, the sample may be treated to lyse the cells, using known lysis buffers, electroporation, etc., with purification and/or amplification such as PCR occurring as needed, as will be appreciated by those in the art. For example, an in vitro transcription with labels covalently attached to the nucleosides is done. Generally, the nucleic acids are labeled with biotin-FITC or PE, or with cy3 or cy5.

In a preferred embodiment, the target sequence is labeled with, for example, a fluorescent, a chemiluminescent, a chemical, or a radioactive signal, to provide a means of detecting the target sequence's specific binding to a probe. The label also can be an enzyme, such as, alkaline phosphatase or horseradish peroxidase, which when provided with an appropriate substrate produces a product that can be detected. Alternatively, the label can be a labeled compound or small molecule, such as an enzyme inhibitor, that binds but is not catalyzed or altered by the enzyme. The label also can be a moiety or compound, such as, an epitope tag or biotin which specifically binds to streptavidin. For the example of biotin, the streptavidin is labeled as described above, thereby, providing a detectable signal for the bound target sequence. As known in the art, unbound labeled streptavidin is removed prior to analysis.

As will be appreciated by those in the art, these assays can be direct hybridization assays or can comprise "sandwich assays", which include the use of multiple probes, as is generally outlined in U.S. Patent Nos. 5,681,702, 5,597,909, 5,545,730, 5,594,117, 5,591,584, 5,571,670, 5,580,731, 5,571,670, 5,591,584, 5,624,802, 5,635,352, 5,594,118, 5,359,100, 5,124,246 and 5,681,697, all of which are hereby incorporated by reference. In this embodiment, in general, the target nucleic acid is prepared

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as outlined above, and then added to the biochip comprising a plurality of nucleic acid probes, under conditions that allow the formation of a hybridization complex.

A variety of hybridization conditions may be used in the present invention, including high, moderate and low stringency conditions as outlined above. The assays are generally run under stringency conditions which allows formation of the label probe hybridization complex only in the presence of target. Stringency can be controlled by altering a step parameter that is a thermodynamic variable, including, but not limited to, temperature, formamide concentration, salt concentration, chaotropic salt concentration pH, organic solvent concentration, etc.

These parameters may also be used to control non-specific binding, as is generally outlined in U.S. Patent No. 5,681,697. Thus it may be desirable to perform certain steps at higher stringency conditions to reduce non-specific binding.

The reactions outlined herein may be accomplished in a variety of ways, as will be appreciated by those in the art. Components of the reaction may be added simultaneously, or sequentially, in any order, with preferred embodiments outlined below. In addition, the reaction may include a variety of other reagents may be included in the assays. These include reagents like salts, buffers, neutral proteins, e.g. albumin, detergents, etc which may be used to facilitate optimal hybridization and detection, and/or reduce non-specific or background interactions. Also reagents that otherwise improve the efficiency of the assay, such as protease inhibitors, nuclease inhibitors, anti-microbial agents, etc., may be used, depending on the sample preparation methods and purity of the target.

Once the assay is run, the data is analyzed to determine the expression levels, and changes in expression levels as between states, of individual genes, forming a gene expression profile.

The screens are done to identify drugs or bioactive agents that modulate the breast cancer phenotype. Specifically, there are several types of screens that can be run. A preferred embodiment is in the screening of candidate agents that can induce or suppress a particular expression profile, thus preferably generating the associated phenotype. That is, candidate agents that can mimic or produce an expression profile in breast cancer similar to the expression profile of normal breast tissue is expected to result in a suppression of the breast cancer phenotype. Thus, in this embodiment, mimicking an expression profile, or changing one profile to another, is the goal.

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In a preferred embodiment, as for the diagnosis and prognosis applications, having identified the differentially expressed genes important in any one state, screens can be run to alter the expression of the genes individually. That is, screening for modulation of regulation of expression of a single gene can be done; that is, rather than try to mimic all or part of an expression profile, screening for regulation of individual genes can be done. Thus, for example, particularly in the case of target genes whose presence or absence is unique between two states, screening is done for modulators of the target gene expression.

In a preferred embodiment, screening is done to alter the biological function of the expression product of the differentially expressed gene. Again, having identified the importance of a gene in a particular state, screening for agents that bind and/or modulate the biological activity of the gene product can be run as is more fully outlined below.

Thus, screening of candidate agents that modulate the breast cancer phenotype either at the gene expression level or the protein level can be done.

In addition screens can be done for novel genes that are induced in response to a candidate agent. After identifying a candidate agent based upon its ability to suppress a breast cancer expression pattern leading to a normal expression pattern, or modulate a single breast cancer gene expression profile so as to mimic the expression of the gene from normal tissue, a screen as described above can be performed to identify genes that are specifically modulated in response to the agent. Comparing expression profiles between normal tissue and agent treated breast cancer tissue reveals genes that are not expressed in normal tissue or breast cancer tissue, but are expressed in agent treated tissue. These agent specific sequences can be identified and used by any of the methods described herein for breast cancer genes or proteins. In particular these sequences and the proteins they encode find use in marking or identifying agent treated cells. In addition, antibodies can be raised against the agent induced proteins and used to target novel therapeutics to the treated breast cancer tissue sample.

Thus, in one embodiment, a candidate agent is administered to a population of breast cancer cells, that thus has an associated breast cancer expression profile. By "administration" or "contacting" herein is meant that the candidate agent is added to the cells in such a manner as to allow the agent to act upon the cell, whether by uptake and intracellular action, or by action at the cell surface. In some embodiments, nucleic acid encoding a proteinaceous candidate agent (i.e. a peptide) may be

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put into a viral construct such as a retroviral construct and added to the cell, such that expression of the peptide agent is accomplished; see PCT US97/01019, hereby expressly incorporated by reference.

Once the candidate agent has been administered to the cells, the cells can be washed if desired and are allowed to incubate under preferably physiological conditions for some period of time. The cells are then harvested and a new gene expression profile is generated, as outlined herein.

Thus, for example, breast cancer tissue may be screened for agents that reduce or suppress the breast cancer phenotype. A change in at least one gene of the expression profile indicates that the agent has an effect on breast cancer activity. By defining such a signature for the breast cancer phenotype, screens for new drugs that alter the phenotype can be devised. With this approach, the drug target need not be known and need not be represented in the original expression screening platform, nor does the level of transcript for the target protein need to change.

In a preferred embodiment, as outlined above, screens may be done on individual genes and gene products (proteins). That is, having identified a particular differentially expressed gene as important in a particular state, screening of modulators of either the expression of the gene or the gene product itself can be done. The gene products of differentially expressed genes are sometimes referred to herein as "breast cancer proteins", "breast cancer modulating proteins" "BCP" or a "BCMP". In one embodiment, BCMP is termed BCH1. In one embodiment, BCH1 can be identified as described for identifying breast cancer proteins herein. In a preferred embodiment, BCH1 is depicted in Figure 34. The BCMP may be a fragment, or alternatively, be the full length protein to the fragment shown herein. Preferably, the BCMP is a fragment of approximately 14 to 24 amino acids long. More preferably the fragment is a soluble fragment.

In other preferred embodiments herein, the breast cancer protein or nucleic acid encoding a breast cancer protein may be selected from any sequence provided in the figures including those wherein the accession numbers are provided. Preferred sequences are in Figure 10, more preferably Figure 11 and most preferably Figure 12. Preferred sequences are also selected from the group consisting of BCH1, BCA2, BCJ7, BCN1, BCN5, BCO2, BCQ5, BCR2, BCR3, BCQ8, BCN2, BCX3, BCX2 and BCY3. Most preferably, the sequence is selected from the group consisting of BCH1, BCA2, BCJ7, BCN1, BCN5, BCO2, BCQ5, BCR2, BCX2 and BCY3. It is understood that any protein can also be

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selected from any subset of this group of proteins, for example, BCH1 can be selected from the whole group, or from the subset of BCH1 and BCN1.

In one embodiment the breast cancer proteins are conjugated to an immunogenic agent as discussed herein. In one embodiment the breast cancer protein is conjugated to BSA.

Thus, in a preferred embodiment, screening for modulators of expression of specific genes can be done. This will be done as outlined above, but in general the expression of only one or a few genes are evaluated.

In a preferred embodiment, screens are designed to first find candidate agents that can bind to differentially expressed proteins, and then these agents may be used in assays that evaluate the ability of the candidate agent to modulate differentially expressed activity. Thus, as will be appreciated by those in the art, there are a number of different assays which may be run; binding assays and activity assays.

In a preferred embodiment, binding assays are done. In general, purified or isolated gene product is used; that is, the gene products of one or more differentially expressed nucleic acids are made. In general, this is done as is known in the art. For example, antibodies are generated to the protein gene products, and standard immunoassays are run to determine the amount of protein present.

Alternatively, cells comprising the breast cancer proteins can be used in the assays.

Thus, in a preferred embodiment, the methods comprise combining a breast cancer protein and a candidate bioactive agent, and determining the binding of the candidate agent to the breast cancer protein. Preferred embodiments utilize the human breast cancer protein, although other mammalian proteins may also be used, for example for the development of animal models of human disease. In some embodiments, as outlined herein, variant or derivative breast cancer proteins may be used.

Generally, in a preferred embodiment of the methods herein, the breast cancer protein or the candidate agent is non-diffusably bound to an insoluble support having isolated sample receiving areas (e.g. a microtiter plate, an array, etc.). The insoluble supports may be made of any composition to which the compositions can be bound, is readily separated from soluble material, and is otherwise compatible with the overall method of screening. The surface of such supports may be solid or porous and of any convenient shape. Examples of suitable insoluble supports include microtiter plates,

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arrays, membranes and beads. These are typically made of glass, plastic (e.g., polystyrene), polysaccharides, nylon or nitrocellulose, teflonTM, etc. Microtiter plates and arrays are especially convenient because a large number of assays can be carried out simultaneously, using small amounts of reagents and samples. The particular manner of binding of the composition is not crucial so long as it is compatible with the reagents and overall methods of the invention, maintains the activity of the composition and is nondiffusable. Preferred methods of binding include the use of antibodies (which do not sterically block either the ligand binding site or activation sequence when the protein is bound to the support), direct binding to "sticky" or ionic supports, chemical crosslinking, the synthesis of the protein or agent on the surface, etc. Following binding of the protein or agent, excess unbound material is removed by washing. The sample receiving areas may then be blocked through incubation with bovine serum albumin (BSA), casein or other innocuous protein or other moiety.

In a preferred embodiment, the breast cancer protein is bound to the support, and a candidate bioactive agent is added to the assay. Alternatively, the candidate agent is bound to the support and the breast cancer protein is added. Novel binding agents include specific antibodies, non-natural binding agents identified in screens of chemical libraries, peptide analogs, etc. Of particular interest are screening assays for agents that have a low toxicity for human cells. A wide variety of assays may be used for this purpose, including labeled in vitro protein-protein binding assays, electrophoretic mobility shift assays, immunoassays for protein binding, functional assays (phosphorylation assays, etc.) and the like.

The determination of the binding of the candidate bioactive agent to the breast cancer protein may be done in a number of ways. In a preferred embodiment, the candidate bioactive agent is labelled, and binding determined directly. For example, this may be done by attaching all or a portion of the breast cancer protein to a solid support, adding a labelled candidate agent (for example a fluorescent label), washing off excess reagent, and determining whether the label is present on the solid support. Various blocking and washing steps may be utilized as is known in the art.

By "labeled" herein is meant that the compound is either directly or indirectly labeled with a label which provides a detectable signal, e.g. radioisotope, fluorescers, enzyme, antibodies, particles such as magnetic particles, chemiluminescers, or specific binding molecules, etc. Specific binding molecules include pairs, such as biotin and streptavidin, digoxin and antidigoxin etc. For the specific binding members, the complementary member would normally be labeled with a molecule which provides for

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detection, in accordance with known procedures, as outlined above. The label can directly or indirectly provide a detectable signal.

In some embodiments, only one of the components is labeled. For example, the proteins (or proteinaceous candidate agents) may be labeled at tyrosine positions using ¹²⁵I, or with fluorophores. Alternatively, more than one component may be labeled with different labels; using ¹²⁵I for the proteins, for example, and a fluorophor for the candidate agents.

In a preferred embodiment, the binding of the candidate bioactive agent is determined through the use of competitive binding assays. In this embodiment, the competitor is a binding moiety known to bind to the target molecule (i.e. breast cancer), such as an antibody, peptide, binding partner, ligand, etc. Under certain circumstances, there may be competitive binding as between the bioactive agent and the binding moiety, with the binding moiety displacing the bioactive agent.

In one embodiment, the candidate bioactive agent is labeled. Either the candidate bioactive agent, or the competitor, or both, is added first to the protein for a time sufficient to allow binding, if present. Incubations may be performed at any temperature which facilitates optimal activity, typically between 4 and 40°C. Incubation periods are selected for optimum activity, but may also be optimized to facilitate rapid high through put screening. Typically between 0.1 and 1 hour will be sufficient. Excess reagent is generally removed or washed away. The second component is then added, and the presence or absence of the labeled component is followed, to indicate binding.

In a preferred embodiment, the competitor is added first, followed by the candidate bioactive agent. Displacement of the competitor is an indication that the candidate bioactive agent is binding to the breast cancer protein and thus is capable of binding to, and potentially modulating, the activity of the breast cancer protein. In this embodiment, either component can be labeled. Thus, for example, if the competitor is labeled, the presence of label in the wash solution indicates displacement by the agent. Alternatively, if the candidate bioactive agent is labeled, the presence of the label on the support indicates displacement.

In an alternative embodiment, the candidate bioactive agent is added first, with incubation and washing, followed by the competitor. The absence of binding by the competitor may indicate that the bioactive agent is bound to the breast cancer protein with a higher affinity. Thus, if the candidate

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bioactive agent is labeled, the presence of the label on the support, coupled with a lack of competitor binding, may indicate that the candidate agent is capable of binding to the breast cancer protein.

In a preferred embodiment, the methods comprise differential screening to identity bioactive agents that are capable of modulating the activity of the breast cancer proteins. In this embodiment, the methods comprise combining a breast cancer protein and a competitor in a first sample. A second sample comprises a candidate bioactive agent, a breast cancer protein and a competitor. The binding of the competitor is determined for both samples, and a change, or difference in binding between the two samples indicates the presence of an agent capable of binding to the breast cancer protein and potentially modulating its activity. That is, if the binding of the competitor is different in the second sample relative to the first sample, the agent is capable of binding to the breast cancer protein.

Alternatively, a preferred embodiment utilizes differential screening to identify drug candidates that bind to the native breast cancer protein, but cannot bind to modified breast cancer proteins. The structure of the breast cancer protein may be modeled, and used in rational drug design to synthesize agents that interact with that site. Drug candidates that affect breast cancer bioactivity are also identified by screening drugs for the ability to either enhance or reduce the activity of the protein.

Positive controls and negative controls may be used in the assays. Preferably all control and test samples are performed in at least triplicate to obtain statistically significant results. Incubation of all samples is for a time sufficient for the binding of the agent to the protein. Following incubation, all samples are washed free of non-specifically bound material and the amount of bound, generally labeled agent determined. For example, where a radiolabel is employed, the samples may be counted in a scintillation counter to determine the amount of bound compound.

A variety of other reagents may be included in the screening assays. These include reagents like salts, neutral proteins, e.g. albumin, detergents, etc which may be used to facilitate optimal protein-protein binding and/or reduce non-specific or background interactions. Also reagents that otherwise improve the efficiency of the assay, such as protease inhibitors, nuclease inhibitors, anti-microbial agents, etc., may be used. The mixture of components may be added in any order that provides for the requisite binding.

Screening for agents that modulate the activity of breast cancer proteins may also be done. In a preferred embodiment, methods for screening for a bioactive agent capable of modulating the activity

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of breast cancer proteins comprise the steps of adding a candidate bioactive agent to a sample of breast cancer proteins, as above, and determining an alteration in the biological activity of breast cancer proteins. "Modulating the activity of breast cancer" includes an increase in activity, a decrease in activity, or a change in the type or kind of activity present. Thus, in this embodiment, the candidate agent should both bind to breast cancer proteins (although this may not be necessary), and alter its biological or biochemical activity as defined herein. The methods include both in vitro screening methods, as are generally outlined above, and in vivo screening of cells for alterations in the presence, distribution, activity or amount of breast cancer proteins.

Thus, in this embodiment, the methods comprise combining a breast cancer sample and a candidate bioactive agent, and evaluating the effect on breast cancer activity. By "breast cancer activity" or grammatical equivalents herein is meant one of the breast cancer's biological activities, including, but not limited to, cell division, preferably in breast tissue, cell proliferation, tumor growth, transformation of cells. In one embodiment, breast cancer activity include activation of breast cancer protein, BCH1, for example, or a substrate thereof by the breast cancer protein. An inhibitor of breast cancer activity is the inhibition of any one or more breast cancer activities.

In a preferred embodiment, the activity of the breast cancer protein is increased; in another preferred embodiment, the activity of the breast cancer protein is decreased. Thus, bioactive agents that are antagonists are preferred in some embodiments, and bioactive agents that are agonists may be preferred in other embodiments.

In a preferred embodiment, the invention provides methods for screening for bioactive agents capable of modulating the activity of a breast cancer protein. The methods comprise adding a candidate bioactive agent, as defined above, to a cell comprising breast cancer proteins. Preferred cell types include almost any cell. The cells contain a recombinant nucleic acid that encodes a breast cancer protein. In a preferred embodiment, a library of candidate agents are tested on a plurality of cells.

In one aspect, the assays are evaluated in the presence or absence or previous or subsequent exposure of physiological signals, for example hormones, antibodies, peptides, antigens, cytokines, growth factors, action potentials, pharmacological agents including chemotherapeutics, radiation, carcinogenics, or other cells (i.e. cell-cell contacts). In another example, the determinations are determined at different stages of the cell cycle process.

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In this way, bioactive agents are identified. Compounds with pharmacological activity are able to enhance or interfere with the activity of the breast cancer protein. In one embodiment, "breast cancer protein activity" as used herein includes at least one of the following: breast cancer activity, binding to a breast cancer protein, activation of a breast cancer protein or activation of substrates of a breast cancer protein by a breast cancer protein.

In one embodiment, a method of inhibiting breast cancer cell division is provided. The method comprises administration of a breast cancer inhibitor.

Neutralizing means that activity of a protein is either inhibited or counter acted against so as to have substantially no effect on the cell.

In another embodiment, a method of inhibiting tumor growth is provided. The method comprises administration of a breast cancer inhibitor.

In a further embodiment, methods of treating cells or individuals with cancer are provided. The method comprises administration of a breast cancer inhibitor.

In one embodiment, a breast cancer inhibitor is an antibody as discussed above. In another embodiment, the breast cancer inhibitor is an antisense molecule. Antisense molecules as used herein include antisense or sense oligonucleotides comprising a singe-stranded nucleic acid sequence (either RNA or DNA) capable of binding to target mRNA (sense) or DNA (antisense) sequences for breast cancer molecules. A preferred antisense molecule is for BCH1 or for a ligand or activator thereof. Antisense or sense oligonucleotides, according to the present invention, comprise a fragment generally at least about 14 nucleotides, preferably from about 14 to 30 nucleotides. The ability to derive an antisense or a sense oligonucleotide, based upon a cDNA sequence encoding a given protein is described in, for example, Stein and Cohen (Cancer Res. 48:2659, 1988) and van der Krol et al. (BioTechniques 6:958, 1988).

Antisense molecules may be introduced into a cell containing the target nucleotide sequence by formation of a conjugate with a ligand binding molecule, as described in WO 91/04753. Suitable ligand binding molecules include, but are not limited to, cell surface receptors, growth factors, other cytokines, or other ligands that bind to cell surface receptors. Preferably, conjugation of the ligand binding molecule does not substantially interfere with the ability of the ligand binding molecule to bind

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to its corresponding molecule or receptor, or block entry of the sense or antisense oligonucleotide or its conjugated version into the cell. Alternatively, a sense or an antisense oligonucleotide may be introduced into a cell containing the target nucleic acid sequence by formation of an oligonucleotide-lipid complex, as described in WO 90/10448. It is understood that the use of antisense molecules or knock out and knock in models may also be used in screening assays as discussed above, in addition to methods of treatment.

The compounds having the desired pharmacological activity may be administered in a physiologically acceptable carrier to a host, as previously described. The agents may be administered in a variety of ways, orally, parenterally e.g., subcutaneously, intraperitoneally, intravascularly, etc. Depending upon the manner of introduction, the compounds may be formulated in a variety of ways. The concentration of therapeutically active compound in the formulation may vary from about 0.1-100 wt.%. The agents may be administered alone or in combination with other treatments, i.e., radiation.

The pharmaceutical compositions can be prepared in various forms, such as granules, tablets, pills, suppositories, capsules, suspensions, salves, lotions and the like. Pharmaceutical grade organic or inorganic carriers and/or diluents suitable for oral and topical use can be used to make up compositions containing the therapeutically-active compounds. Diluents known to the art include aqueous media, vegetable and animal oils and fats. Stabilizing agents, wetting and emulsifying agents, salts for varying the osmotic pressure or buffers for securing an adequate pH value, and skin penetration enhancers can be used as auxiliary agents.

Without being bound by theory, it appears that the various breast cancer sequences are important in breast cancer. Accordingly, disorders based on mutant or variant breast cancer genes may be determined. In one embodiment, the invention provides methods for identifying cells containing variant breast cancer genes comprising determining all or part of the sequence of at least one endogeneous breast cancer genes in a cell. As will be appreciated by those in the art, this may be done using any number of sequencing techniques. In a preferred embodiment, the invention provides methods of identifying the breast cancer genotype of an individual comprising determining all or part of the sequence of at least one breast cancer gene of the individual. This is generally done in at least one tissue of the individual, and may include the evaluation of a number of tissues or different samples of the same tissue. The method may include comparing the sequence of the sequenced breast cancer gene to a known breast cancer gene, i.e. a wild-type gene.

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The sequence of all or part of the breast cancer gene can then be compared to the sequence of a known breast cancer gene to determine if any differences exist. This can be done using any number of known homology programs, such as Bestfit, etc. In a preferred embodiment, the presence of a a difference in the sequence between the breast cancer gene of the patient and the known breast cancer gene is indicative of a disease state or a propensity for a disease state, as outlined herein.

In a preferred embodiment, the breast cancer genes are used as probes to determine the number of copies of the breast cancer gene in the genome.

In another preferred embodiment breast cancer genes are used as probed to determine the chromosomal localization of the breast cancer genes. Information such as chromosomal localization finds use in providing a diagnosis or prognosis in particular when chromosomal abnormalities such as translocations, and the like are identified in breast cancer gene loci.

Thus, in one embodiment, methods of modulating breast cancer in cells or organisms are provided. In one embodiment, the methods comprise administering to a cell an anti-breast cancer antibody that reduces or eliminates the biological activity of an endogenous breast cancer protein. Alternatively, the methods comprise administering to a cell or organism a recombinant nucleic acid encoding a breast cancer protein. As will be appreciated by those in the art, this may be accomplished in any number of ways. In a preferred embodiment, for example when the breast cancer sequence is down-regulated in breast cancer, the activity of the breast cancer gene is increased by increasing the amount of breast cancer in the cell, for example by overexpressing the endogenous breast cancer or by administering a gene encoding the breast cancer sequence, using known gene-therapy techniques, for example. In a preferred embodiment, the gene therapy techniques include the incorporation of the exogenous gene using enhanced homologous recombination (EHR), for example as described in PCT/US93/03868, hereby incorporated by reference in its entirety. Alternatively, for example when the breast cancer sequence is up-regulated in breast cancer, the activity of the endogeneous breast cancer gene is decreased, for example by the administration of a breast cancer antisense nucleic acid.

In one embodiment, the breast cancer proteins of the present invention may be used to generate polyclonal and monoclonal antibodies to breast cancer proteins, which are useful as described herein. Similarly, the breast cancer proteins can be coupled, using standard technology, to affinity chromatography columns. These columns may then be used to purify breast cancer antibodies. In a preferred embodiment, the antibodies are generated to epitopes unique to a breast cancer protein;

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that is, the antibodies show little or no cross-reactivity to other proteins. These antibodies find use in a number of applications. For example, the breast cancer antibodies may be coupled to standard affinity chromatography columns and used to purify breast cancer proteins. The antibodies may also be used as blocking polypeptides, as outlined above, since they will specifically bind to the breast cancer protein.

In one embodiment, a therapeutically effective dose of a breast cancer or modulator thereof is administered to a patient. By "therapeutically effective dose" herein is meant a dose that produces the effects for which it is administered. The exact dose will depend on the purpose of the treatment, and will be ascertainable by one skilled in the art using known techniques. As is known in the art, adjustments for protein degradation, systemic versus localized delivery, and rate of new protease synthesis, as well as the age, body weight, general health, sex, diet, time of administration, drug interaction and the severity of the condition may be necessary, and will be ascertainable with routine experimentation by those skilled in the art.

A "patient" for the purposes of the present invention includes both humans and other animals, particularly mammals, and organisms. Thus the methods are applicable to both human therapy and veterinary applications. In the preferred embodiment the patient is a mammal, and in the most preferred embodiment the patient is human.

The administration of the breast cancer proteins and modulators of the present invention can be done in a variety of ways as discussed above, including, but not limited to, orally, subcutaneously, intravenously, intranasally, transdermally, intraperitoneally, intramuscularly, intrapulmonary, vaginally, rectally, or intraocularly. In some instances, for example, in the treatment of wounds and inflammation, the breast cancer proteins and modulators may be directly applied as a solution or spray.

The pharmaceutical compositions of the present invention comprise a breast cancer protein in a form suitable for administration to a patient. In the preferred embodiment, the pharmaceutical compositions are in a water soluble form, such as being present as pharmaceutically acceptable salts, which is meant to include both acid and base addition salts. "Pharmaceutically acceptable acid addition salt" refers to those salts that retain the biological effectiveness of the free bases and that are not biologically or otherwise undesirable, formed with inorganic acids such as hydrochloric acid, hydrobromic acid, sulfuric acid, nitric acid, phosphoric acid and the like, and organic acids such as

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acetic acid, propionic acid, glycolic acid, pyruvic acid, oxalic acid, maleic acid, malonic acid, succinic acid, fumaric acid, tartaric acid, citric acid, benzoic acid, cinnamic acid, mandelic acid, methanesulfonic acid, ethanesulfonic acid, p-toluenesulfonic acid, salicylic acid and the like.

"Pharmaceutically acceptable base addition salts" include those derived from inorganic bases such as sodium, potassium, lithium, ammonium, calcium, magnesium, iron, zinc, copper, manganese, aluminum salts and the like. Particularly preferred are the ammonium, potassium, sodium, calcium, and magnesium salts. Salts derived from pharmaceutically acceptable organic non-toxic bases include salts of primary, secondary, and tertiary amines, substituted amines including naturally occurring substituted amines, cyclic amines and basic ion exchange resins, such as isopropylamine, trimethylamine, diethylamine, triethylamine, tripropylamine, and ethanolamine.

The pharmaceutical compositions may also include one or more of the following: carrier proteins such as serum albumin; buffers; fillers such as microcrystalline cellulose, lactose, corn and other starches; binding agents; sweeteners and other flavoring agents; coloring agents; and polyethylene glycol. Additives are well known in the art, and are used in a variety of formulations.

In a preferred embodiment, breast cancer proteins and modulators are administered as therapeutic agents, and can be formulated as outlined above. Similarly, breast cancer genes (including both the full-length sequence, partial sequences, or regulatory sequences of the breast cancer coding regions) can be administered in gene therapy applications, as is known in the art. These breast cancer genes can include antisense applications, either as gene therapy (i.e. for incorporation into the genome) or as antisense compositions, as will be appreciated by those in the art.

In a preferred embodiment, breast cancer genes are administered as DNA vaccines, either single genes or combinations of breast cancer genes. Naked DNA vaccines are generally known in the art. Brower, Nature Biotechnology, 16:1304-1305 (1998).

In one embodiment, breast cancer genes of the present invention are used as DNA vaccines. Methods for the use of genes as DNA vaccines are well known to one of ordinary skill in the art, and include placing a breast cancer gene or portion of a breast cancer gene under the control of a promoter for expression in a breast cancer patient. The breast cancer gene used for DNA vaccines can encode full-length breast cancer proteins, but more preferably encodes portions of the breast cancer proteins including peptides derived from the breast cancer protein. In a preferred embodiment a patient is immunized with a DNA vaccine comprising a plurality of nucleotide sequences derived

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from a breast cancer gene. Similarly, it is possible to immunize a patient with a plurality of breast cancer genes or portions thereof as defined herein. Without being bound by theory, expression of the polypeptide encoded by the DNA vaccine, cytotoxic T-cells, helper T-cells and antibodies are induced which recognize and destroy or eliminate cells expressing breast cancer proteins.

In a preferred embodiment, the DNA vaccines include a gene encoding an adjuvant molecule with the DNA vaccine. Such adjuvant molecules include cytokines that increase the immunogenic response to the breast cancer polypeptide encoded by the DNA vaccine. Additional or alternative adjuvants are known to those of ordinary skill in the art and find use in the invention.

In another preferred embodiment breast cancer genes find use in generating animal models of breast cancer. As is appreciated by one of ordinary skill in the art, when the breast cancer gene identified is repressed or diminished in breast cancer tissue, gene therapy technology wherein antisense RNA directed to the breast cancer gene will also diminish or repress expression of the gene. An animal generated as such serves as an animal model of breast cancer that finds use in screening bioactive drug candidates. Similarly, gene knockout technology, for example as a result of homologous recombination with an appropriate gene targeting vector, will result in the absence of the breast cancer protein. When desired, tissue-specific expression or knockout of the breast cancer protein may be necessary.

It is also possible that the breast cancer protein is overexpressed in breast cancer. As such, transgenic animals can be generated that overexpress the breast cancer protein. Depending on the desired expression level, promoters of various strengths can be employed to express the transgene. Also, the number of copies of the integrated transgene can be determined and compared for a determination of the expression level of the transgene. Animals generated by such methods find use as animal models of breast cancer and are additionally useful in screening for bioactive molecules to treat breast cancer.

It is understood that the examples described herein in no way serve to limit the true scope of this invention, but rather are presented for illustrative purposes. All references and sequences of accession numbers cited herein are incorporated by reference in their entirety.

EXAMPLE

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Expression studies were performed herein on both the protein and nucleic acid level. Results are provided in the figures.

An experiment was performed to examine the correlation (if any) between BCH1 protein and estrogen receptor (ER) expression and localization. Antibodies against BCH1 protein were obtained by immunization in rabbits with 15-mer synthetic peptides corresponding to regions of BCH1. Anti-ER antibodies were purchased commercially. Approximately 50 breast carcinoma tissue specimens (5 micron serial sections; paraffin-embedded, formalin-fixed) were analyzed by immunohistochemistry. Of these, ~40% were positive for BCH1 expression; ~35% were positive for ER. Protein levels were scored from 1 to 5 by immunohistochemistry, with 5 representing the highest expression. Co-expression of BCH1 and ER is shown in Figure 37.

Of the ER+ tumors (n=108), about one-half have predominantly nuclear localizations of ER, one-half predominantly cytoplasmic. Interestingly, while the BCH1+/ER+ tumors were no more prevalent than by chance, without exception tumors with high levels of BCH1 (IHC score = 4 or 5) showed exclusively localization of ER to the cytoplasm, with exclusion from the nucleus (see Figures 38 and 39) (n=16, p<5 x 10^{-7}), whereas low expression of BCH1 had no correlation with ER localization and was often associated with nuclear ER (see Figures 40 and 41).

Currently, ER expression is an important clinical prognostic marker for breast cancer. Because signaling through ER to activate most estrogen-responsive genes is believed to require translocation of activated ER to the nucleus, high expression of BCH1 is predicted to correlate with functionally-negative ER in ER+ tumors. Since ER- correlates with poor prognosis, BCH1 may also be a valuable prognostic marker, as it may also correlate with poor prognosis, or to no-responsiveness to antiestrogen therapies (e.g., tamoxifen), since approximately 50% of the ER+ patients do not respond to tamoxifen. Tamoxifen works against the effects of estrogen on these cells. It is often called an "antiestrogen." As a treatment for breast cancer, the drug slows or stops the growth of cancer cells that are already present in the body. As adjuvant therapy, tamoxifen has been shown to help prevent the original breast cancer from returning and also prevent the development of new cancers in the opposite breast.

Recent publications report that ER can signal through the Ras/Raf/MAP kinase. Since many of the proteins in this signaling cascade reside in the cytoplasm, increased BCH1 expression may correlate with, or even cause, ER to remain in the cytoplasm, thereby altering the signaling of ER through the

MAP kinase pathway. This may be an explanation for the subset of ER+ patients who do not respond to tamoxifen.

Furthermore, BCH1 may have a casual role in ER translocation to the nucleus, by either directly or indirectly preventing nuclear translocation, thereby eliminating or altering responsiveness of the cell to estrogens or their analogs.

- 1. A method of screening drug candidates comprising:
 - a) providing a cell that expresses an expression profile gene which encodes a protein selected from the group consisting of BCH1, BCA2, BCJ7, BCN1, BCN5, BCO2, BCQ5, BCR2, BCX2 and BCY3 or a fragment thereof;
 - b) adding a drug candidate to said cell; and
 - c) determining the effect of said drug candidate on the expression of said expression profile gene.
- 2. A method according to claim 1 wherein said determining comprises comparing the level of expression in the absence of said drug candidate to the level of expression in the presence of said drug candidate, wherein the concentration of said drug candidate can vary when present, and wherein said comparison can occur after addition or removal of the drug candidate.
- 3. A method according to claim 1 wherein the expression of said profile gene is decreased as a result of the introduction of the drug candidate.
- 4. A method of screening for a bioactive agent capable of binding to a breast cancer modulator protein (BCMP), wherein said BCMP is BCH1 or a fragment thereof, said method comprising combining said BCMP and a candidate bioactive agent, and determining the binding of said candidate agent to said BCMP.
- 5. A method for screening for a bioactive agent capable of modulating the activity of a breast cancer modulator protein (BCMP), wherein said BCMP is BCH1 or a fragment thereof, said method comprising combining said BCMP and a candidate bioactive agent, and determining the effect of said candidate agent on the bioactivity of said BCMP.
 - 6. A method of evaluating the effect of a candidate breast cancer drug comprising:
 - a) administering said drug to a patient;
 - b) removing a cell sample from said patient; and
 - c) determining the expression profile of said cell.

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- 7. A method according to claim 6 further comprising comparing said expression profile to an expression profile of a healthy individual.
- 8. A biochip comprising a nucleic acid segment encoding BCH1 or a fragment thereof, wherein said biochip comprises fewer than 1000 nucleic acid probes.
- 5 9. A method of diagnosing breast cancer comprising:
 - a) determining the expression of a gene encoding BCH1 or a fragment thereof in a first tissue type of a first individual; and
 - b) comparing said expression of said gene from a second normal tissue type from said first individual or a second unaffected individual;

wherein a difference in said expression indicates that the first individual has breast cancer.

- 10. An antibody which specifically binds to BCH1, or a fragment thereof.
- 11. The antibody of Claim 10 wherein said fragment is BCH1p1 or BCH1p2.
- 12. The antibody of Claim 10, wherein said antibody is a monoclonal antibody.
- 13. The antibody of Claim 10, wherein said antibody is a humanized antibody.
- 14. The antibody of Claim 10, wherein said antibody is an antibody fragment.
 - 15. A method for screening for a bioactive agent capable of interfering with the binding of a breast cancer modulator protein (BCMP) or a fragment thereof and an antibody which binds to said BCMP or fragment thereof, said method comprising:
 - a) combining a BCMP or fragment thereof, a candidate bioactive agent and an antibody which binds to said BCMP or fragment thereof; and
 - b) determining the binding of said BCMP or fragment thereof and said antibody.
 - 16. A method for inhibiting breast cancer, said method comprising administering to a cell a composition comprising an antibody to BCH1 or a fragment thereof.
 - 17. The method of Claim 16 wherein said cell is a cell of an individual.

- 18. The method of Claim 17 wherein said individual has cancer.
- 19. The method of Claim 16 wherein said fragment is selected from the group consisting of BCH1p1 and BCH1p2.
- 20. The method of Claim 16 wherein said antibody is a humanized antibody.
- 5 21. The method of Claim 16 wherein said antibody is an antibody fragment.
 - 22. A method for inhibiting breast cancer in a cell, wherein said method comprises administering to a cell a composition comprising antisense molecules to BCH1.
 - 23. A peptide consisting essentially of BCH1p1.
 - 24. A composition comprising the peptide of Claim 23.
 - 25. A peptide consisting essentially of BCH1p2.
 - 26. A composition comprising the peptide of Claim 25.
 - 27. A method of eliciting an immune response in an individual, said method comprising administering to said individual a composition comprising BCH1 or a fragment thereof.
- 28. A method of eliciting an immune response in an individual, said method comprising administering to said individual a composition comprising a nucleic acid comprising a sequence encoding BCH1 or a fragment thereof.
 - 29. A composition capable of eliciting an immune response in an individual, said composition comprising BCH1 or a fragment thereof and a pharmaceutically acceptable carrier.
- 30. A composition capable of eliciting an immune response in an individual, said composition comprising a nucleic acid comprising a sequence encoding BCH1 or a fragment thereof and a pharmaceutically acceptable carrier.

- 31. A method of treating an individual for breast cancer comprising administering to said individual an inhibitor of BCH1.
- 32. The method of Claim 31 wherein said inhibitor is an antibody.
- 33. The method of Claim 31 wherein said individual is non-responsive to an anti-estrogen and is positive for estrogen receptor.
 - 34. The method of Claim 33 wherein said method further comprises administering an antiestrogen.
 - 35. A method for determining the prognosis of an individual with breast cancer comprising determining the level of BCH1 in a sample, wherein a high level of BCH1 indicates a poor prognosis.
 - 36. A method for determining whether an individual with breast cancer will be non-responsive to anti-estrogen therapies comprising determining the level of BCH1 wherein a high level of BCH1 indicates that an individual will be non-responsive.
 - 37. A method of neutralizing the effect of a BCH1, or a fragment thereof, comprising contacting an agent specific for said protein with said protein in an amount sufficient to effect neutralization.

ABSTRACT

Described herein are methods that can be used for diagnosis and prognosis of breast cancer. Also described herein are methods that can be used to screen candidate bioactive agents for the ability to modulate breast cancer. Additionally, methods and molecular targets (genes and their products) for therapeutic intervention in breast and other cancers are described.

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Come Name	ESTs ESTS EST	Insulin-like growth factor binding protein 6 EST - HG1428-HT1428	EST - HG2157-HT2227 EST - HG2841-HT2969	Fatty acid binding protein 4 adipocyte SERUM AMYLOID A PROTEIN PRECURSOR	Activating transcription factor 3	Alcohol dehydrogenase 1 (class I) alpha polypeptide EST - M21305	Phospholipase A2 group IIA (platelets synovial	Human endogenous retroviral protease mRNA	Cholesteryl ester transfer protein plasma EST - U22961 Homo sapiens protein kinase C-binding protein	EST - U88902_cds1_f PLASMA RETINOL-BINDING PROTEIN Interleukin 6 (B cell stimulatory factor 2) SERUM AMYLOID A PROTEIN PRECURSOR Tetranectin (plasminogen-binding protein) TrkB {alternatively spliced} [human brain mRNA	H.sapiens mRNA for hHKb1 protein Homo sapiens MIP-1 delta mRNA complete cds EST - AA081995	ESTS ESTS ESTS ESTS ESTS ESTS	ESTs
,	Affymetrix ID	RC_R16733 R16733 RC_AA079072 AA079072 HG1428-HT1428 TIGR - HG1428-HT1428	-	- 5 5	, H	L49169 L49169 M12963 M12963		M25079 M25079 M27826 M27826	M30185 M30185 U22961 U22961 U48251 U48251	U88902 cds1 f U88902 X00129 X00129 X04602 X04602 X51441 X51441 X64559 X64559 X75958 X75958	X9/544_Custo_t X9/544 X99142 X99142 Z49269 Z49269 A A 081995 A A 081995	. 4 4 4 .	AA285284 AA285284 AA310850 AA310850

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AA418143 AA425719 AA427379 AA452705 AB002328 AF000575	M12272 M26315 M64936	M81349 M84526 N73185 N79674 N88827 N91071	R69417 AA009764 AA017254 AA019300 AA016280 AA115253 AA1179338 AA223237	AA234308 AA251772 AA279673 AA416947 AA426584 AA434113 AA443303 AA443303 AA449471 AA449471
AA418143 AA425719 AA427379 AA452705 AB002328 AF000575_s	M12272_s I M26315_cds2_s I M64936_i I	M81349 M84526 N73185 N79674_s N88827 N91071_s R21149	R69417 RC_AA009764 RC_AA017254 RC_AA019300 RC_AA015253 RC_AA115253 RC_AA115338 RC_AA179338 RC_AA179338	RC_AA234308 RC_AA251772 RC_AA219673 RC_AA416947 RC_AA446947 RC_AA443303_ RC_AA443303_ RC_AA443471 RC_AA449471 RC_AA449471

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ESTs Weakly similar to ORF YOR173w ESTs ESTs Homo sapiens transmembrane protein mRNA	Homo sapiens Kruppel-like zinc finger protein	EST	ES 18 EST - RC_D59420 EST - T87593	Human MEK5 mRNA complete cds	Fullial Will ob Illary Complete cus	ESTs Human mRNA for rah GDI alnha complete cds	ESTs	ESTs Weakly similar to D2030.9 [C.elegans] EST - W38002 s	Interleukin 6 (B cell stimulatory factor 2)	Cholinergic receptor nicotinic delta polypeptide ESTs	ESTs Weakly similar to C27H6.5 [C.elegans] ESTs	ESTs	ESTS	SISE TSE	ESTs Weakly similar to HYPOTHETICAL 41.9 KD	ESTs	ESTs	EST - RC_AA075124	EST - RC_AA079079	EST - RC: AAU/9120	ESI - RC_AAU830/U_S	ESTS ESTS	
RC_AA478487 AA478487 RC_AA491001_f AA491001 RC_AA620446 AA620446 RC_AA621131 AA621131 RC_AA621414_ AA621414	RC_AA621680 AA621680	, F 6	RC_D56989_i_D56989 RC_D59420_D59420 _T87593_T87593		- •	W26097 W26097 W28390		W28931 W28931 W38002 s W38002	တ	X55019_s X55019 RC AA011576 AA011576			•	RC_AA037388 AA037388	RC_AA047229 AA047229	RC AA059473 AA059473			RC_AA079079 AA079079		•	RC_AA121820 AA121820 RC_AA126583 AA126583	

FIGURE 1 (CONT.)

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ESTs ESTs ESTs ESTs ESTs ESTs ESTs ESTs	ESTs Weakly similar to HYPOTHETICAL 46.1 KD	ESTs ESTs Homo sapiens clone 23770 mRNA sequence ESTs ESTs ESTs ESTs ESTs ESTs ESTs ESTs
RC_AA131571 AA131571 RC_AA219555_ AA219555 RC_HA219555_ AA219555 RC_H05645_ H05645 RC_H10761_H10761 RC_H18299_H18299 RC_H51276_H5299 RC_H51276_H58394 RC_H58934_H58934 RC_H69547_H69547	RC_N49409 N49409	RC_N62889_s N62889 RC_N66951 N66951 RC_R00144 R00144 RC_R07324 R07324 RC_R3146 R3146 RC_R3146 R3146 RC_R42333 R42333 RC_R43977 R43977 RC_R43977 R43977 RC_R43977 R43977 RC_R66992 R66992 RC_R77302 R77302 RC_R77302 R77302 RC_T01803 T03803 RC_T58756 T58756 RC_T79768 T79768 RC_T91047 T91047 RC_W04657 W04657 RC_W86195 W86195 RC_AA435850 AA443800 RC_AA445800 AA443800 RC_AA445806 AA456968 RC_AA456968 AA456968 RC_AA456968 AA456968

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10.0	10.0	10.0 10.0 10.0 10.0 10.0 10.0 10.0	10.0 10.0 10.0 10.0 10.0	10.0	10.0	10.0	10.0	10.0
EST Homo sapiens intermediate conductance calciumactivated potassium channel (hKCa4) mRNA	TYROSINE-PROTEIN KINASE RECEPTOR TIE-	ESTs Weakly similar to line-1 protein ORF2 ESTs ESTs ESTs ESTs ESTs EST Human G0S2 protein gene complete cds EST - RC_W38051 EST MYO-INOSITOL-1(OR 4)-	ESTs EST EST ESTs ESTS EST Human apM1 mRNA for GS3109 (novel adipose	ESTs Moderately similar to nuclear autoantigen	Homo sapiens oligodendrocyte-specific protein	P55-C-FOS PROTO-ONCOGENE PROTEIN ESTs Highly similar to HYPOTHETICAL 38.2 KD PROTEIN IN BEM2-SPT2 INTERGENIC	ESTs Cytochrome P450 subfamily XXI (steroid 21-	H.sapiens KHK mRNA for ketohexokinase clone
RC_C21161 C21161 RC_D20860 D20860	RC_H02848_s H02848	RC_N20468 N20468 RC_N39426 N39426 RC_N49285 f N49285 RC_N50034 N50034 RC_N5972 N65972 RC_N70907 N70907 RC_T47418 T47418 RC_T48075 f T48075 RC_T52813 s T52813 RC_W38051 W38051 RC_W38051 W38051 RC_W3523 W73523 RC_AA223746 f AA223746	RC_AA431337 AA431337 RC_AA447555 AA447555 RC_AA458945 AA458945 RC_AA485421_f AA485421 RC_AA621529_f AA621529 RC_H15814_s H15814	RC_H90310_r H90310	RC_N22392 N22392	RC_N23730_s N23730 RC_N50809 N50809	RC_R48732_s_R48732 RC_T47089_s_T47089	RC_T61256_s T61256

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10.0 10.0 10.0 10.0 5.0	5.0 5.0 5.0 5.0	5.0 5.0 5.0	5.0 5.0 5.0 5.0	5.0 5.0	5.0 5.0 5.0 5.0	5.0 5.0	5.0 5.0	5.0	5.0
ESTs ESTs Homo sapiens mRNA for perilipin complete cds Homo sapiens transmembrane protein mRNA	EST - HG2147-HT2217 r EST - HG2796-HT2904 EST - HG3236-HT3413 f EST - HG537-HT537 DIHYDROPRYRIDINE-SENSITIVE L-TYPE	CELL SURFACE GLYCOPROTEIN A15 Pregnancy-associated plasma protein A Lipocalin 1 (protein migrating faster than albumin	GRO2 oncogene Insulin-like growth factor binding protein 6 Human G0S2 protein gene complete cds Prostaglandin E receptor 3 (subtype EP3)	EST - U32674 Homo sapiens skeletal muscle LIM-protein FHL1	Glutamate receptor metabotropic 4 Alcohol dehydrogenase 2 (class I) beta polypeptide Tenascin R (restrictin janusin) Hemoglobin alpha 1 ESTs Weakly similar to ZINC FINGER PROTEIN	Homo sapiens clone 23718 mRNA sequence Homo sapiens mRNA for GNAS1 protein (IMAGE	EST - AA082561_s Homo sapiens secreted frizzled related protein	ESTs Weakly similar to HYPOTHETICAL 39.7 KD	EST - AA191072
RC_T94447_s T94447 RC_T98199_f T98199 RC_W72887 W72887 RC_W94688 W94688 AF000959 AF000959	HG2147- TIGR - HG2147-HT2217 HG2796-HT2904 TIGR - HG2796-HT2904 HG3236- TIGR - HG3236-HT3413 HG537-HT537 TIGR - HG537-HT537 L07738 L07738	L10373 L10373 L13197 L13197 L14927 L14927	M57731 M57731 M62402 M62402 M72885_ma1 M72885 S68874 S68874	U32674 U32674 U60115 U60115	U92457 U92457 X03350 X03350 X98085 X98085 Z84721 cds2 Z84721 AA044622 AA044622	AA059327_r AA059327 AA062932 AA062932	AA082561_s AA082561 AA093348 AA093348	AA166651 AA166651	AA191072 AA191072

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5.0 5.0 5.0 5.0 5.0 5.0 5.0 5.0 5.0 5.0 5.0 5.0 5.0 5.0 5.0 5.0 5.0 ESTs ESTs Amine oxidase copper containing 2 (retina-specific) EST - F15201 ESTs Highly similar to GLYCINE-RICH ESTs Weakly similar to L8083.1 gene product Human tyrosyl-tRNA synthetase mRNA complete ESTs Highly similar to OVARIAN GRANULOSA protein (pseudo)gene an unknown putative gene a Human clone 23803 mRNA partial cds ESTs Moderately similar to serine protease ESTs Weakly similar to APK1 antigen [H.sapiens] Human mRNA for KIAA0363 gene partial cds POU domain class 1 transcription factor 1 (Pit1 ESTs Weakly similar to cytoplasmic dynein light Laminin alpha 3 (nicein (150kD) kalinin (165kD) Human DNA sequence from clone 71L16 on ESTs Weakly similar to HYPOTHETICAL Homo sapiens secreted frizzled related protein Transcription factor 3 (E2A immunoglobulin chromosome Xp11. Contains a probable Zinc Finger ESTs Weakly similar to eukaryotic initiation factor AA461426_r AA461426 AB002361 AB002361 AA448946_r AA448946 AA278194 AA278194 AA278194 AA278194 AA291786_s AA291786 AA402109 AA402109 AA416829 AA416829 AA418214 AA418214 AA210757 AA210757 AA232121 r AA232121 AA247434 AA247434 AA400044 AA400044 AA402971_s AA402971 AA422123_i AA422123 AF001900 AF001900 H30778 L34155 D10216_s D10216 N40774 N40774 D88213 D88213 F15201 C16161 s C16161 C17282 C17282 N39361 N39361 D31381 D31381 L34155 F15201 H30778

5.0	5.0	5.0	5.0	5.0	5.0	5.0	2.0	5.0	5.0	5.0	5.0	5.0 5.0	0.4	0.4	0.0	0.0	5.0	5.0	5.0	5.0	5.0	5.0	5.0	2.0	5.0	5.0	5.0	5.0	5.0	5.0 0.0	5.0	5.0	0.0	0.0	0.0
EST	Homo sapiens exportin t mRNA complete cds	ESTs	ESTs Moderately similar to !!!! ALU SUBFAMILY	H.sapiens mRNA for phosphoinositide 3-kinase	ESTs	Acyl-Coenzyme A dehydrogenase very long chain	ESTs	ESTS	ESTs	ESTS	ESIS	ESTs Highly similar to LIGATIN [H.sapiens]	2117 2100	210d	Homo sapiens mKNA for KIAAU652 protein	ESIS	ESIS	ESTS	ESTs Weakly similar to potassium-dependent	Early growth response 2 (Krox-20 (Drosophila)	ESTS	ESTS	Homo sapiens chromosome 19 cosmid R27216	Human mRNA for KIAA0124 gene partial cds	ESTS	ESTS	ESTs	ESTs	ESTs	ESTS	ESTS	Homo sapiens mRNA for doublecortin	Human LMP1 associated protein mRNA complete	ESIS	Homo sapiens killer cell receptor (KIR103) mRNA
N99542 N99542		R24011 R24011	R25944 f R25944	R68735 R68735			RC AA007153 AA007153		-	٠,	~	~	RC_AA1518/2 AA1518/2	RC_AA215643 AA215643		RC_AA402613 AA402613	RC_AA405449 AA405449	RC_AA422146 AA422146	RC_AA427627 AA427627	DC A A 446027 A A 446027		KC_AA433944 AA433944 pC_AA63939 AA633939	•		•	RC_AA487576 AA487576	RC_AA489499 AA489499	-	-	RC_AA608802 AA608802	RC_AA609785 AA609785	RC_AA621430 AA621430		U51704 U51704	U73394_f U73394

5.0	5.0	5.0 5.0 5.0 5.0 5.0 5.0 5.0	5.0 5.0 5.0 5.0 5.0	8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8 8	0.0 0.0 0.0 0.0 0.0 0.0 0.0 0.0 0.0
Human tissue inhibitor of metalloproteinase 4	Human macrophage-derived chemokine precursor	ESTS ESTS ESTS ESTS MAP KINASE PHOSPHATASE-1 MONOCYTE CHEMOTACTIC PROTEIN 3 ESTS ESTS ESTS	EST - RC_AA070500 Homo sapiens macrophage receptor MARCO ESTs ESTs	ESTs Moderately similar to !!!! ALU SÜBFAMILY ESTs ESTs EST - RC_H52172 ESTs ESTs ESTs ESTs ESTS ESTS	ESTs Weakly similar to No definition line found ESTs Highly similar to LATENT TRANSFORMING GROWTH FACTOR BETA ESTS ESTS ESTS ESTS ESTS ESTS ESTS
U76456 U76456	U83171 U83171		RC_AA063174 AA063174 RC_AA070500 AA070500 RC_AA074885 AA074885 RC_AA099820 AA099820	RC_AA207059 AA207059 RC_AA207059 AA207059 RC_H41280 H41280 RC_H52172 H52172 RC_H58222 H58222 RC_N52176 N52176 RC_N6516 N66116 RC_N67583 N67583	RC_N73988 N73988 RC_N92239 N92239 RC_R26065 R26065 RC_R43035 R43035 RC_R51898 R51898 RC_R84968 R84968 RC_R96306 R96306 RC_R98491 R98491 RC_R98491 R98491

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RC_T54659 T54659 RC_T57857 T57857 RC_T89160_f T89160 RC_T91185 T91185 RC_Z39975 Z39975 RC_Z41480 Z41480 RC_AA020736 AA020736	RC_AA412707 AA412707 RC_AA428964 AA428964 RC_AA463902 AA463902 RC_AA488080 AA488080 RC_AA488839 AA488839 RC_F10528_f F10528	RC_H90434 H90434 RC_H97387_s H95958 RC_H97387_s H97387 RC_N23399 N23399 RC_N35978 N35978 RC_N59860_s N59860 RC_N69476 N69476 RC_N69476 N69476 RC_N90744 N90744 RC_R95132 R95132 RC_T41144_s T41144 RC_T777892 T77892 RC_T77892 T77892 RC_W74257 W74257

ESTs 5.0			/TLS 5.0	_								oiens] 5.0			465_f 5.0		NI E:						ESTs 5.0		ESTs 3.3	ESTs 3.3		ESTs 3.3		tein 6 3.3	ES IS 5.3
			RNA-BINDING PROTEIN FUS/TLS						Homo sapiens mRNA for KIAA0538 protein partial	T mondiodrody LISE		ESTs Weakly similar to ADAM 20 [H.sapiens]	AXL receptor tyrosine kinase	EST - RC_H59063	EST - RC_H83465_f	ESTs Highly similar to HYPOTHETICAL 64	GTP-BINDING PROTEIN C02F5.3 IN	Glutathione S-transferase M5	EST - RC_R01398			VON WILLEBKAND FACTOR FRECORSOR FRE		Homo sapiens STAT-induced STAT inhibitor-2					EST - RC_N53145_f	Pregnancy-specific beta-1 glycoprotein 6	
89182M 89182M Od	RC_W/8108 W/6108	KC_W8/555 W8/555		DC A A 283018 A A 383018	٠.	•	RC AA405763 AA405763	RC_AA421483 AA421483	-	•	RC_AA459658 AA459658	RC_AA6091122 AA6091122 RC_AA609214 AA609214	_	ان 	_	RC N27118 N27118	ı	RC N56898 s N56898	RC R01398 R01398	RC R44214 r R44214	RC_R71403_f R71403	RC_T61561_s T61561	RC_T94357_f_T94357		TSSSAT TSSSAT	DC WA4723 WA4733	DC 4 4401564 A 4401564		- :-	 RC_R02384 R02384	RC_W73194 W73194

3333 Tetranectin (plasminogen-binding protein)
ESTs
ESTs
ESTs
ESTT- YEL024w/RIP1

RC_W73889_s W73889 RC_W92278 W92278 RC_R48540_s R48540 YEL024w/RIP1

FIGURE 2

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Ratio breast v. tumor		10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0
Rati		ESTS	ESTs	EST	ESTs	Insulin-like growth factor binding protein	EST - HG1428-HT1428	Human G0S3 mRNA complete cds	SERUM AMYLOID A PROTEIN	D component of complement (adipsin)	ESTs	Homo sapiens Kruppel-like zinc finger	ESTs	ESTs	ESTs	P55-C-FOS PROTO-ONCOGENE	Homo sapiens mRNA for perilipin	EST - HG2157-HT2227	EST - HG2841-HT2969	Fatty acid binding protein 4 adipocyte	SERUM AMYLOID A PROTEIN	Activating transcription factor 3	Alcohol dehydrogenase 1 (class I) alpha	EST - M21305	Phospholipase A2 group IIA (platelets	Hemoglobin beta	Human endogenous retroviral protease	Cholesteryl ester transfer protein plasma	EST - U22961	Homo sapiens protein kinase C-binding	EST - U88902_cds1_f	PLASMA RETINOL-BINDING	Interleukin 6 (B cell stimulatory factor 2)	Tetranectin (plasminogen-binding	TrkB {alternatively spliced} [human	H.sapiens DMA DMB HLA-Z1 IPP2	H.sapiens mRNA for hHKb1 protein	Homo sapiens MIP-1 delta mRNA	EST - AA081995	ESTs
	Accession	AA025277	H49425	N51657	R16733		TIGR - HG1428-	L49169	X51441	M84526	R69417	AA621680	AA071193	N66951	R33146	N23730	W94688	TIGR - HG2157-	TIGR - HG2841-	J02874	J03474	L19871	M12963	M21305	M22430	M25079	M27826	M30185	U22961	U48251	U88902	X00129	X04602	X64559	X75958	X87344	X99142	Z49269	AA081995	AA090439
	Attymetrix ID	RC AA025277	RC H49425	RC_N\$1657	RC_R16733	RC AA079072 s	HG1428-HT1428	L49169	X51441	M84526	R69417	RC AA621680	RC_AA071193	RC N66951	RC_R33146	RC N23730 s	RC W94688	HG2157-HT2227	HG2841-HT2969	102874	303474	L19871	M12963	M21305	M22430	M25079	M27826	M30185	U22961	U48251	U88902 cds1 f	\bar{x} 001 \bar{z}	X04602	X64559	X75958	X87344 cds10 r	_X99142	Z49269	AA081995	AA090439

				s 10.0		•		e 10.0	1 10.0											, , ,			10.0 10.0		_				,	•					_ `		10.0	
ESTs	ESTs	ESTs Highly similar to ISOCITRATE	ESTS	ESTS	ESTs	ESTS	FSTs Weakly similar to D2045.9	Human mRNA for KIAA0330 gene	Homo sapiens clone 17.11	Alcohol dehydrogenase 3 (class I) gamma	CD8 antigen alpha polypeptide (p32)	Homo sapiens retinoic acid-inducible	SERUM AMYLOID A-4 PROTEIN	EST	ESTs	ESTs	ESTs	ESTs Highly similar to RAB GDP	ESTS	ESTs	ESTS	ESIS	ESTS	ESIS		ES Is Moderately summar to removines-	H sanjens mRNA for HES1 protein	Homo sapiens mRNA for HsGAK	ESTS	ESTs	ESTs	ESTS	ESIS	ESTS	ESTS	ESTS	ESIS	ESTs Weakly similar to ORF YORI/3w
AA137107	A A 203296	AA285284	A A 310850	A A 418143	A A 425710	A A A 77379	202024AA	AR452/03	A E000575	M12272	M26315	M64936	M81349	N73185	N79674	N88827	N91071	R21149	AA009764	AA017254	AA019300	AA026280	AA115253	AA128617	AA179338	AA223237	A A 251772	AA279673	AA411443	AA416947	AA426584	AA434113	AA443303	AA446005	AA449471	AA451877	AA478487	AA491001
A A 137107	A A 203296	A A 785784	A A 210850	AA310630	AA416145	AA425/19	AA421319	AA452/05	AB002528	Ar000373_s	2 Caba 215.CM	M64936 i	M81349	N73185	N79674 s	N88827	N91071 s	R21149	RC AA009764		RC_AA019300	RC_AA026280	RC_AA115253	RC_AA128617	RC_AA179338	RC_AA223237	RC_AA234308	DC_A A 279673	RC_AA411443	RC_AA416947	RC_AA426584	RC_AA434113	RC AA443303 s	RC AA446005	RC_AA449471	RC_AA451877	RC_AA478487	RC AA491001_f

10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	10.0	5.0	5.0	5.0	5.0	5.0	5.0
ESTs	Homo sapiens transmembrane protein	EST	ESTs	ESTs	EST - RC_D59420	EST - T87593	Human MEK5 mRNA complete cds	Human Wnt10B mRNA complete cds	ESTs	ESTs	Human mRNA for rab GDI alpha	ESTs	ESTs Weakly similar to D2030.9	Homo sapiens secreted frizzled related	Homo sapiens mRNA for KIAA0652	ESTs	ESTs	ESTs	Apolipoprotein D
AA620446	AA621414 AA621414	D25786	D26989	D26989	D59420	T87593	Ú25265	U81787	W19098	W26097	W28390	W28548	W28931	AF001900	AA402268	AA608802	F01449	AA034378	AA456975
RC_AA620446	RC_AA621131 RC_AA621414_s	RC D25786	RC D56989 f	RC_D56989_i	RC D59420	T87593	U25265	U81787	W19098	W26097	W28390	W28548	W28931	AF001900	RC AA402268	RC_AA608802	RC F01449 f	RC AA034378 F	RC_AA456975_s

FIGURE 3

ratio breast vs tumor Homo sapiens MIP-1 delta mRNA complete cds ESTs Human G0S3 mRNA complete cds
SERUM AMYLOID A PROTEIN PRECURSOR
ESTs Tetranectin (plasminogen-binding protein) Insulin-like growth factor binding protein 6 Cholesteryl ester transfer protein plasma Gene Name EST - U22961 ESTs Accession R33146 M30185 U22961 X64559 Z49269 AA427379 AA079072 L49169 X51441 N66951 Affymetrix ID RC_AA079072_s RC_N66951 RC_R33146 M30185 Z49269 AA427379 X51441 X64559 L49169 U22961

Ratio tumor v. breast 135.3 121.4 107.3 81.8 75.3 72.9 61.5 57.1 53.9 53.2 52.0 46.8 45.7 38.9 33.6 32.4 31.3 31.1	28.1 28.5 25.5 25.1 23.9 23.8 22.8 20.9 19.4 17.7 17.1
ESTs EST-RC_AA453638 EST - RC_AA461510 Collagen type XI alpha 1 EST - RC_N27351 H.sapiens mRNA for Sm protein F Human focal adhesion kinase (FAK) mRNA complete cds ESTs Weakly similar to zinc-finger protein Zn72D ESTs ESTs ESTs ESTS ARYLAMINE N-ACETYLTRANSFERASE ESTS ESTS ESTS ESTS ESTS ESTS ESTS E	EST - RC_AA211831 ESTS Inhibin beta A (activin A activin AB alpha polypeptide) ESTS ESTS Human TFIID subunits TAF20 and TAF15 mRNA ESTS ESTS ESTS ESTS ESTS ESTS ESTS EST
Acession T79956 AA453640 AA453638 AA461322 AA461310 R67275 AA461510 R67275 AA461310 AA453479 AA453479 AA453479 AA453479 AA453641 AA621202 AA453641 AA609955 AA453641	AA211831 AA412090 N27159 AA421289 T16687 R65763 AA487987 H99309 R97063 AA232940 AA463189 AA421171 AA251875 AA651875 AA651875 AA651875 AA651875 AA651875 AA651875
Affymetrix ID RC T79956 RC AA453640 RC AA453638 RC AA461322 RC AA461322 RC AA461510 RC AA461510 RC AA461510 RC AA486737 RC AAA486737 RC AAA486737 RC AAA486737 RC AAA486737 RC AAA53479 RC AAA53479 RC AAA53479 RC AAA53479 RC AAA632294 RC AAA53641 RC AAA53641 RC AAA53641 RC AAA53641	RC_AA211831 RC_AA211831 RC_AA412090 RC_N27159_s RC_AA421289 RC_AA487987 RC_AA487171 RC_AA4871875 RC_AA4871875 RC_AA4871875 RC_AA63189 RC_AA63189 RC_AA63189 RC_AA63189 RC_AA63189

17.0	16.8	16.3	16.2	1,51	16.1	15.9	15.8	15.8	15.7	15.6	15.4	13.2	14.9	14.7	14.6	14.6	14.6	14.4	14.4	14.4	14.4	14.3	14.3	14.2	14.1	14.0	14.0	13.9	13.9	13.9	13.7	13.5	13.5	13.4	13.4
ESTs H.sapiens mRNA for hHKb1 protein	EST - RC_AA211158	Human 4E-binding protein 1 mRNA complete cds	ESTS	C.I. Coll. College Col	ES1s Highly similar to thyroid disease hypothetical FST - RC AA 196721	ESTS	Homo sapiens clone 23967 unknown mRNA partial cds	Inhibin beta A (activin A activin AB alpha polypeptide)	TRANSCOBALAMIN I PRECURSOR	ESTs Moderately similar to !!!! ALU SUBFAMILY SP	ESTS	EDIS	ESTs Weakly similar to B0334.4 [C.elegans]	Human fibroblast activation protein mRNA complete cds	ESTs	ESTs	Interferon regulatory factor 5	ESTs Moderately similar to PTTG gene product	ESTs	ESTs	Carboxypeptidase B1 (tissue)	ESTs Highly similar to HYPOTHETICAL 21.5 KD	Human effector cell protease receptor-1 (EPR-1) gene	ESTs	Chromogranin A (parathyroid secretory protein 1)	ESTs Moderately similar to 25E8.1 [D.melanogaster]	ESTs Weakly similar to TH1 protein [D.melanogaster]	Androgen receptor (dihydrotestosterone receptor testicular	ESTs	ESTs Weakly similar to hypothetical protein 1 [H.sapiens]	ENIS	Homo sapiens histone macroH2A1.2 mRNA complete cds	ESTS Moderately similar to 111G6.8 [C.elegans]	ES 18 WEARLY SHILLIAL TO 22-FAZAYOU AUTOMINISKII	Homo sapiens KIAA0440 mRNA partial cds
AA232508 AA024659	AA488191 AA211158	AA290674	AA481883	AA190/08	H83527	D51177	T25875	X57579	105068	AA487233	AA479969	AA191404	AA262969	AA436611	R51309	AA461297	AA250843	AA430032	AA280679	AA412029	M81057	R07976	U75285	R46627	AA461559	AA092129	AA436893	M23263	AA465345	AA486538	D20379	AA076138	W60486	AA0430/4	F01444
RC_AA232508 RC_AA024659	RC_AA488191 RC_AA211158	RC_AA290674	RC_AA481883	KC_AA196/68	H83527_s	DS DS1172	RC_T25875	X57579	305068	RC AA487233	RC_AA479969	RC_AA191404	RC_AA262969	RC AA436611	RC R51309	RC AA461297	RC_AA250843	RC_AA430032	RC_AA280679	RC_AA412029	M81057	RC R07976	U75285 ma1	RC R46627	RC AA461559	AA092129 f	RC AA436893	_ M23263	RC AA465345	RC_AA486538	RC_D20379	RC_AA076138	RC_W60486	KC_AA045074	RC_F01444_f

13.3 13.3 13.1 13.1	13.0 12.8 12.7 12.5 12.5 12.4	12.3 12.2 12.2 12.2 12.2 12.2	12.1 12.1 11.8 11.5 11.4 11.4	11.3 11.3 11.0 10.9 10.9
ESTs Homo sapiens mRNA for SCGF-beta complete cds Human mRNA for KIAA0007 gene partial cds ESTs Moderately similar to !!!! ALU SUBFAMILY SC EST - RC_AA412065 ESTs	Homo sapiens regulator of G-protein signalling 12 ESTs ESTs EST Homo sapiens mRNA for A+U-rich element RNA binding ESTs ESTS	ESTs Weakly similar to !!!! ALU SUBFAMILY SX Homo sapiens ES/130 mRNA complete cds ESTs ESTs ESTs ESTs	Fibroblast growth factor receptor 2 (bacteria-expressed Glutamine-fructose-6-phosphate transaminase Human Hep27 protein mRNA complete cds ESTs UBIQUITIN-LIKE PROTEIN GDX ESTs ESTs ESTs ESTs ESTs ESTs	Homo sapiens chromosome 9 P1 clone 11659 ESTs Weakly similar to extracellular protein [H.sapiens] EST ESTs Weakly similar to transmembrane protein [H.sapiens] EST Homo sapiens U4/U6 small nuclear ribonucleoprotein EST
AA401334 T78922 D60354 AA431350 AA412065 AA412065	AA431738 R61740 R54950 AA405488 AA418749 AA0377285 AA233796 AA233796	AA463874 AA463874 AA461528 AA099404 AA443985 AA214305	AA220223 AA478571 U31875 AA253217 AA470074 AA236010 D82307 AA430002	R22952 AA179298 W56363 AA44054 AA281733 AA452601 AA035630 AA235117 AA279418
AA401334 RC_T78922_s RC_D60354_s RC_AA431350 RC_AA412065 RC_AA406635	RC_AA431738 RC_R61740_f RC_R61740_f RC_AA405488 RC_AA418749 AA037285 RC_AA219305	RC_AA452245 RC_AA41276 RC_AA463874 RC_AA461528 RC_AA499404 RC_AA443985	AA220223 RC_AA478571 U31875 RC_AA253217 RC_AA470074 RC_AA236010 J03589 D82307 RC_AA430002	RC_A22952_s RC_AA179298 RC_AA449232 RC_AA44054 RC_AA44054 RC_AA481733 RC_AA482601 RC_AA482601 RC_AA482601 RC_AA482601 RC_AA482601 RC_AA482601 RC_AA482601

10.8 10.7 10.7	10.6 10.5 10.5	10.4	10.3 10.3	10.2	10.1 10.0 10.0	9.9 9.9	x 8 6 6	7.0 7.0 7.0	9.7 9.6 9.6	9.6 6.9 9.5	,	<i>!</i>
ESTs EST EST PST	ESTs Weakly similar to MOESIN/EZRIN/RADIXIN ESTs FSTs	Human splicesomal protein (SAP 61) mRNA complete cds GANGLIOSIDE GM2 ACTIVATOR PRECURSOR FST	ESTs ESTs Highly similar to RAS-RELATED PROTEIN RAB-	ESTs Highly similar to ZINC FINGER PROTEIN 85 ESTs Weakly similar to K02B2.3 gene product [C.elegans]	ESTs ESTs Weakly similar to trabecular meshwork inducible ESTs Weakly similar to No definition line found	H.sapiens DAP-3 mRNA EST	ESTS ESTS ESTS	Spermidine synthase ESTs FSTs Weakly similar to R01H10.6 [C.elegans]	PROTEASOME COMPONENT C13 PRECURSOR EST EST EST-RC_T97341	ESTs Highly similar to OVOSTATIN PRECURSOR ESTs Highly similar to HYPOTHETICAL 23.1 KD ESTs	ESTS ESTS Highly similar to PUTATIVE ATP-DEPENDENT ESTS ESTS ESTS ESTS ESTS	1 1 2 2
AA432069 AA453630 W44657	AA405098 AA411425	AA599259 X62078	AA253170 AA459347 AA251430 AA470156	AA280609	AA449832 AA427898 AA609867	R49198 AA465158	AA112396 AA207015 R06986	M34338 AA228030 A A 447982	AA44,7982 Z14982 AA176247 T97341	W26392 AA143190 AA452578	W92113 AA282914 AA461476 AA258057 W87751 N21678 AA26111	11170707
RC_AA432069 RC_AA453630 RC_W44657	RC_R40431 RC_AA405098 RC_AA411425	RC_AA59259 RC_AA599259 X62078	RC_AA253170 RC_AA459347 RC_AA251430	RC_AA4/0130 RC_T64933_f RC_AA281290 RC_AA280609	RC_AA449832 RC_AA427898 RC_AA609867	RC_R49198_i RC_AA465158	RC_AA112396 RC_AA207015 RC_R06986_f	M34338 RC_AA228030 PC_AA447082	RC_AA44,7962 Z14982_ma1 RC_AA176247 RC_T97341	W26392 RC_AA143190 RC_AA452578	RC_W92/13 RC_AA282914 RC_AA461476 RC_AA258057 RC_W87751 RC_N21678	KC_AA202111

9.3	9.3	9.3	9.2	9.2	9.2	9.1	9.1	9.1	9.1 9.1	9.0	9.0	0.6	0.6	y. o	0.0	× × ×	9 6	%·\	8.7	%. <u>\</u>	8.6	0.0	o «	200	% 	 	4.0	† v	. o «	; eq	, c.		8.3	8.2	8.2	8.1	8.1	8.1	
EST	ESTs	ESTs	ESTs	ESTs Highly similar to TRYPSINOGEN ANIONIC	Human mRNA for KIAA0101 gene complete cds	ESTs	Human CENP-F kinetochore protein mRNA complete cds	H.sapiens mRNA for disintegrin-metalloprotease (partial)	ESTS	EST	ESTS	Human germline IgD chain gene C-region C-delta-1	EST - HG2981-H1312/	S100 calcium-binding protein A7 (psoriasin 1)	H.sapiens Humig mKINA	Human mRNA for KIAA0324 gene partial cds	ESIS	ESIS	ESIS	MULTIFUNCTIONAL AMINOACYL-TRNA	Homo sapiens clone 23785 mRNA sequence	ESIS	TRANSFORMATION-SENSITIVE PROTEIN 1EF 5SP	CLOH.	Human mRNA for KIAA0314 gene partial cds	Collagen type XI alpha 1	Homo sapiens Shab-related delayed-rectifier K+ channel	FIGHIORIAN	\$155 1015AT OG T92	FOLD IN FIRST	HONE TO THE STATE OF THE ACT AND ACT ACT	True alone 22716 mDNA sequence	FSTS	FSTs Highly similar to HYPOTHETICAL 37.8 KD	Matrix metalloproteinase 2 (gelatinase A collagenase type	Homo saniens mRNA for KIAA0617 protein complete cds	ESTs	ESTs	
0 A A 490020	0500EV	N80716	A A 007344	W73140	D14657	T16308	AA447666	AA243020	AA431478	R38919	R60223	R70379	TIGR - HG2981-	M86757	X72755	AA347209	AA485041	AA443342	AA481281	T96361	H18027	AA608723	M86752	AA457018	AA113011	H96237	AA024835	S85655	9266N	T65004	/4154N	H/2948	AA489510	75035CA A	1050270	H98621	R40177	D60302	
0.000000	KC_AA490929	RC_N/0090	RC_1460/10	DC W73140	D14657	PC T16308 f	RC A 447666	RC_AA243020	RC_AA431478	RC R38919 i	RC_R60223_s	RC_R70379_s	_HG2981-	M86757	X72755	RC AA347209	RC_AA485041	RC_AA443342	RC_AA481281	RC T96361 s	RC_H18027_s	RC AA608723	_ M86752	RC_AA457018	RC_AA113011	RC H96237_s	RC_AA024835	S85655	RC_N99976	RC_T65004	RC_N93197	RC_H72948_s		KC_2399/1_s	KC_AA23005/	0/0000 2 109601 2 d	KC_130021_3	RC_D60302	

8.1	8.0	8.0	8.0	8.0	7.9	7.9	7.9	7.9	7.9	7.9	7.9	7.9	7.9	7.9	7.8	7.8	7.8	7.8	7.8	7.8	7.8	7.7	7.7	7.7	9.7	2.6	7.6	7.6	2.6	9.2	7.5	7.5	7.5	7.5	7.5	7.5	7.5	7.5
Human mariner-like element-containing mRNA clone	ESTs Weakly similar to HYPOTHETICAL 26.1 KD	EST's Highly similar to OSTEOINDUCTIVE FACTOR	ES	RAP1 GTPase activating pro-	ESTS	ESTs	ESTs Weakly similar to LINE-1 REVERSE	Replication protein A (E coli RecA homolog RAD51	EST	ESTs	ESTs	ESTs	ESTs	EST - RC_N53950	ESTs Weakly similar to HYPOTHETICAL 88.1 KD	Homo sapiens Ran binding protein 2 (RanBP2alpha)	EST - RC_AA342084	ESTs	EST	EST - RC_N46435	Human mRNA for KIAA0136 gene partial cds	ESTs	Homo sapiens mRNA for tyrosyl sulfotransferase-2	ESTs	ESTs Highly similar to EPIDERMAL GROWTH	Human melanoma antigen p15 mRNA complete cds	ESTs	EST	Homo sapiens clone 24703 beta-tubulin mRNA complete	EST - HG2981-HT3938	ESTs	Tyrosine 3-monooxygenase/tryptophan 5-monooxygenase	Human mRNA for KIAA0170 gene complete cds	ESTs	EST - W01296	ESTs		Homo sapiens mRNA for osteoblast specific factor 2 (OSF-
AA419225	A A 733545	N39415	AA436370	F01538	AA053319	. N69464	AA458882	N33011	AA421750	AA235009	AA447574	R01634	R47948	N53950	AA150182	AA446486	AA342084	AA417213	AA609170	N46435	N54916	T88814	AA459389	AA463693	H99879	019796	T68871	AA446008	T03306	TIGR - HG2981-	N67119	AA442767	AA454566	AA476937	W01296	AA282074	AA094752	D13666
RC_AA419225	BC 4 4 233545	RC N39415	RC AA436370	RC_F01538_s	RC_AA053319	RC N69464	RC AA458882	RC N33011 s	RC_AA421750	RC_AA235009	RC_AA447574	RC R01634	RC R47948 i	RC N53950	RC AA150182	RC_AA446486	RC_AA342084	RC_AA417213	RC_AA609170	RC N46435	RC_N54916	RC_T88814	RC AA459389	RC_AA463693	RC_H99879	U19796	RC_T68871	RC_AA446008	RC_T03306	HG2981-	RC N67119	RC AA442767	RC_AA454566	RC_AA476937	_ W01296	RC_AA282074	AA094752	D13666

7.4	7.4	7.4	7.4	7.4	7.4	7.4	7.4	7.4	7.4	7.4	7.4	7.4	7.3	7.3	5.7	7.3	 	7.5 5.7	7.7	7:7 7.7	4: L	, L	4. C	7.1	7.1	7.1	7.1	7.1	7.1	7.1	7.1	7.1	7.1	7.1	7.1	7.0	7.0	
ESTs	ESTs	EST	ESTs	EST	ESTS Highly similar to HYPOTHETICAL 28.5 KD	H saniens mRNA for surface glycoprotein	ESTS	Homo saniens mRNA for GalT4 protein	ESTs	FCTs Moderately similar to breast cancer suppressor	Interferon (gamma)-induced cell line protein 10 from	ESTs	ESTs Highly similar to CHROMOSOME	ESTs	ESIS	Human mRNA for histone H1x complete cds	EDIS	ESIS	ESIS	Homo sapiens mRNA for kynurenine 3-monooxygenase	ESI - RC AA42/950	ESTs Weakly sumilar to !!!! ALU SUBFAINIL I	ESTs Moderately similar to !!!! ALU SUBFAMILT SA	[2000] 11 0 4 10 T	ESTs Weakly similar to 104A8.11 [Celegans]	Transfer of the malana arough regulatory protein	H. Sapiens mixara tot metanoma grown regument from ESTs	ESTs	ESTs	ESTs Highly similar to HYPOTHETICAL 52.8 KD	Human mRNA for zinc finger protein complete cds	Lymphotoxin-beta	ESISE	521S 1530CM 7d 753	124547 - 163 1845	SICA Standard Contraction of the	ESTS Weakly similar to similar to T06D8.5 [Celegans]	EDIS WEAKLY SILLILLE TO ACCOUNT LETTERS AND ACCOUNTS.
N67889	002001	A A 112/17	AA412477	AA439392	AA399042	W /3320	AA009470	AA28/001	AA411952	AA410190	X02530	050704	AA283006	AA443794	F13642	AA426372	AA446869	N21321	AA121315	R65593	AA427950	AA088458	AA432130	AA234921	AA310967	AA236177	AA282143	AAZ83003	T10082	C00225	AA258482	AA287870	AA410373	F13694	N29431	N67239	AA449351	240345
0862314 04	KC_N0/009	KC_AA609309	RC_AA412477	RC_AA459392	RC_AA599042	RC_W73520	RC_AA069476	RC_AA287061	RC_AA411952	RC_AA410190	RC_AA486256	08033X	KC_D39489	DC_A A 443794	PC F13642	RC AA426372	BC A A 446869	RC N21321 i	BC 44121315	RC R65593 s	RC AA427950	RC_AA088458	RC_AA432130	RC_AA234921	AA310967_s	RC_AA236177	RC_AA282143	RC_AA283003	RC AA421130	1,10001, JA	DC A A 258487	BC_A A 287870	RC_AA410373	RC F13694 f	RC N29431	RC_N67239	RC_AA449351	RC_Z40345

7.0	7.0	7.0	6.9	6.9	6.9	6.9	8.9	8.9	8.9	6. 8	× •	o.x	۰.۵ د د	ø.0 .0	۰.0 م	Ø 1	6.7	6.7	6.7	6.7	6.7	6.7	6.7	6.7	6.7	6.7	6.6	9.9	9.9	0.0	9.9	9.9	9.9	9.9	9.9	0.0	6.5	. v	0.0
Homo saniens clone 23915 mRNA sequence	ESTs	EST	ESTs	ESTs	ESTs	Receptor protein-tyrosine kinase EDDR1	ESTs	ESTs	ESTs Highly similar to VALYL-TRNA SYNTHETASE	ESTs Highly similar to putative hydrophobic domain in	Human clone 23589 mRNA sequence	ESTS	ESTs Weakly similar to !!!! ALU SUBFAMILY J	H.sapiens mRNA for FAST kinase	ESTs Weakly similar to TRANSFORMATION-	ESI	Homo sapiens mRNA for Efs1 complete cds	Human autoantigen pericentriol material 1 (PCM-1)	Human bumetanide-sensitive Na-K-Cl cotransporter	ESTs	ESTS	Human JTV-1 (JTV-1) mRNA complete cds	Interleukin 6 signal transducer (gp130 oncostatin M	ESTs Moderately similar to unknown protein [H.sapiens]	ESTs	Collagen type V alpha	ESTs	ESTs	ESTS	ESIS	ESTs Highly similar to FK506-BINDING PROTEIN	ESTs Highly similar to SKD3 [M.musculus]	ESTs	ESTs	Nuclear factor of kappa light polypeptide gene enhancer in	ESTS	ESTs Highly similar to COP1 REGULATORY PROTEIN	ESI	Human prolyl 4-hydroxylase alpha (II) subunit mKNA
N34686	A A 310400	08573G	N21204	A A 410441	A A 505093	1148705	AA419461	AA411204	AA496569	AA346385	D51229	AA127818	H18428	F02254	AA416876	AA446966	NS0550	L27841	U30246	AA034069	T92935	U24169	H99935	AA435849	AA127058	M11718	AA283198	R51988	AA505141	D60341	N26904	R40606	T03790	W72455	AA098834	AA421782	AA236384	AA431085	W49521
2027674 04	KC_N34686	NC_AA310439	DO 171707	TO 14/1/04	DC_A4410441		RC A A 419461	RC_AA411204	RC_AA496569	RC_AA346385	RC D51229 f	RC AA127818	RC H18428 s	RC_F02254_s	RC AA416876	RC_AA446966	RC N50550		1130246	RC AA034069	RC T92935	U24169	RC H99935 s	RC AA435849	RC_AA127058	M11718	RC AA283198	RC R51988	RC AA505141	RC D60341	RC_N26904	RC_R40606	RC_T03790	RC W72455	RC AA098834	RC_AA421782	_AA236384	RC AA431085	_ W49521

6.5	6.5	6.4	6.4	6.4	6.4	6.4	6.4	6.3	6.3	6.3	6.3	6.3	6.3	6.3	6.3	6.3	6.3	6.3	6.2	6.2	6.2	6.2	6.2	6.2	6.2	6.2	6.2	6.2	6.1	6.1	6.1	6.1	6.1	6.1	6.1	6.1	6.1	6.1
ESTS	ESTs	MULTIFUNCTIONAL AMINOACYL-TRNA	Human pyridoxal kinase mRNA complete cds	EST	ESTs	ESTs	ESTs	ESTs	Homo sapiens DDX1 gene complete CDS	EST	H.sapiens mRNA for RNA polymerase II subunit	Homo sapiens mRNA for KIAA0549 protein partial cds	ESTs	Homo sapiens COX17 mRNA complete cds	Homo sapiens lamin B receptor homolog TM7SF2	ESTs	Human mRNA for KIAA0050 gene complete cds	ESTs	Human NADH:ubiquinone oxidoreductase subunit B13	ESTs	ESTs	H.sapiens mRNA for MUF1 protein	ESTs	ESTs	Protein-tyrosine kinase 7	Arylsulfatase A	ESTs	Homo sapiens chaperonin containing t-complex	LAMIN B1	Human eukaryotic translation initiation factor (eIF3)	Tubulin gamma polypeptide	ESTs	ESTs Highly similar to GOLIATH PROTEIN [Drosophila	Homo sapiens mRNA for putative RNA helicase 3' end	ESTs	ESTs	ESTs	ESTs
AA446591 R06700	AA100364	X54326	90968N	AA195651	T15991	AA430211	T17119	AA157814	X70649	H57330	D81608	R65826	AA621169	L77701	AA443658	W19662	D20168	N48166	AA024664	AA279943	AA098874	X86018	AA412106	AA621721	U40271	X52150	AA152178	N38959	L37747	U78525	T77733	D20280	W69807	AA405505	AA133199	HS5748	AA479933	AA448349
RC_AA446591 RC_R06700	RC AA100364	X54326	90968N	RC_AA195651	RC T15991	RC AA430211	RC T17119	RC AA157814	X70649	RC H57330	_D81608	RC R65826	RC AA621169	_ L77701	RC AA443658	W19662	RC D20168	RC_N48166	RC AA024664	RC_AA279943	RC_AA098874	X86018	RC AA412106	RC_AA621721	_ U40271	X52150 mal s	RC AAI52178	RC N38959 f	_L37747_s	$0.785\overline{2}5$	RC T77733 s	$R\bar{C}$ D20280	RC W69807	RC AA405505	RC_AA133199	RC H55748	RC AA479933	RC_AA448349

6.1 6.1 6.1	6.1	6.1	6.1	6.1	6.1	9.0	0.0	0.0	0.9	6.0	0.9	5.9	5.9	5.9	5.9	6.0	6.5	5.9	5.9	5.8	5.8	5.8	5.8	×. v	0.0	0.0	×. ×	×. ×	0.0	o. «	. 8. 8.	5.8	5.7
ERGIC-53 PROTEIN PRECURSOR ESTs Weakly similar to line-1 protein ORF2 [H.sapiens] ESTs	ESTs Weakly similar to LINE-1 REVERSE	ESTS	ESTs	ESTS	Human gerruline oligomeric matrix protein (COMP)	ESIS	ESIS	ED LS	Home conjens short chain 13-hydroxvacyl-CoA	ESTs	Collagen type XI alpha 1	ESTs	ESTs	ESTs	ESTs Weakly similar to keratin 8 type II cytoskeletal	\$15a	STOTI STOTI	II. II. Services of the stackment factor B (SAF-B)	Homo sapiens scattord anachment acces 2 (273, 23) EST - HG3748-HT4018	ESTs	ESTs	Cyclin B1	ESTs	ESTs Weakly similar to PROCOLLAGEN ALPHA 1(II)	ESIS	ENIS	ESTS	CATHEPSIN K PRECURSOR	ENIS	Human multidrug resistance-associated protein homolog	ES IS FSTs	Human extracellular matrix protein 1 (ECM1) mRNA	ESTs
AA600257 R99978	AA455551 H55915	AA398740 A A 521080	AA416568	N92593	L32137	AA609277	R33663	AA521103	AA406137	T16660	110000	W38407	N26391	A A 292655	Z41619	AA223730	T88953	NS0/44	W63563	-04/CDII - VOII A A 430673	A A 463740	M25753	AA279292	AA427925	AA287665	AA422007	AA425379	T67463	AA441801	H89987	H94843	C80108NI C805911	N27563
RC_AA600257 RC_R99978	AA455331 RC_H55915	RC_AA398740	RC_AA16568	RC N92593		RC_AA609277	RC_R33663_s	RC_AA521103	RC_AA406137	0/3314	KC_116660	71417 DC W38407	PC_N26491	PC A A 202655	RC_Z41619_s	RC_AA223730	RC_T88953	RC_N50744	RC_W63563_s	<	PC_A4430073	M25753	RC AA279292	RC_AA427925	RC_AA287665	RC_AA422007	RC_AA425379	RC T67463 s	RC AA441801	RC_H89987_s	RC_H94843	RC_N80183 _1165027	RC_N27563

5.7	5.7	5.7	5.7	5.7	5.7	5.7	5.7	5.7	5.7	7.0	7.0	5.7	5.7	5.7	5.6	5.6	5.6	5.6	5.6	5.6	0.0	5.0	5.0	0.0 V	5.6	2.0	5.6	5.6	5.6	5.6	5.6	5.6	5.6	5.6	5.6	
ESTs	Fibronectin 1 Fibronectin 1	Homo sapiens mRNA for follistain-related protein (FRP)	Homo sapiens mRNA for KIAA0585 protein partial cds	SICE NEW YOR OF WORTH IIIV GOTTA AT	FACTOR VIII IN INCIA 24 1 NO LEIN	ESTS	ESTs	ESTs	Homo sapiens clone 24522 mRNA sequence	ESTS	ESI EST	FST	EST	EST - HG2981-HT3125	ESTs	EST - U91327	ESTs	Homo sapiens KIAA0431 mRNA partial cds	ESTs	ESTs	ESIS	ESTs Moderately similar to TYKi protein [M.musculus]	Human mRNA for KIAA0124 gene partial cds	H.sapiens DAP-1 mRNA	E21	ES1S FRT	Stock	Guanine nucleonde oniquing protein O protein EST	ESTS	ECTs Weakly similar to T12D8.i [C.elegans]	ESTS Weard Stimmer to the ESTS	Homo saniens clone 23736 mRNA sequence	Human fatty acid amide hydrolase mRNA complete cds	ESTs	Integral transmembrane protein l	
F09058	R02572	AA442763 A A 149624	AA459945	R09166	M34677	AA478794	W30943	A A 454567	U56402	AA443251	AA207105	AA609473	H54430	K83829	110K - HG2961- A A 232956	1191377	N51590	AA406169	AA147884	AA453987	AA040154	T23528	D50914	X76105	AA149754	AA397919	AA398212	AA416986	AA435936	195051	AA116093	777174V	A A 435742	AA452842	AA482269	
RC F09058	RC_R02572	RC_AA442763 pC_AA149624		RC_R09166	M34677	RC_AA478794	W30943	RC_AA192334 nC_AA64662	KC_AA434302 1156402	AA443251	RC_AA207105	RC_AA609473	RC_H54430	RC_R85829	-1867DH	KC_AA232930	12150 721500 a	RC_A406169	RC_AA147884	RC_AA453987	RC_AA040154	RC T23528	D50914	X76105	RC AA149754	RC_AA397919	RC_AA398212	RC_AA416986	RC_AA435936	RC_T95057_f	AA116095	AA4//214	KC_AA398204	DC A 4457847	RC_AA482269	

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0.0 4 4 4 2 6 8 1 0	A A 436819	ENIS	0.0
N93797	N93797	ESTs	5.6
2C A 443602	AA443602	ESTS	5.5
RC_A A 609996	AA609996	ESTs Highly similar to Surf-4 protein [M.musculus]	C. C.
RC_AA075200	AA075200	Homo sapiens Chromosome 16 BAC clone C1198/SK-A-	5. 5. 5. 5.
RC_AA195517	AA195517	ESTs Weakly similar to !!!! ALU SUBFAMILL I 3	ر از د
_ X02874	X02874	E. S. S. Shiniciase E. (25.) oligoadenyiate symmetase E. S.	, v.
RC_AA085589	AA085589	ESTS Highly similar to 1 KANOLA 1101N MALLA 1101N	5.5
RC_H99500	H99500	Homo sapiens mixive for formstantification process. ESTs	5.5
RC_R43883	K43883	December of the 2-oxoglutarate 4-dioxygenase (proline	5.5
M24486	M24480	FIOCOIDAGEII-PLOMIC Z-OAGEMMAN CONTRACTOR CO	5.5
RC_AA115535	AA115555	ESTs	5.5
RC_AA280840	AA280840	ESTs	5.5
RC_N22015	C1022NI	ESTs	5.5
RC_AA021182	AA021102	EST	5.5
KC_N21032	1109278	Human fibroblast activation protein mRNA complete cds	5.5
4	A A 251973	ESTs	5.4
KC AA2319/3	H75933	Laminin receptor (2H5 epitope)	5.4
KC_f1/3935_1	T81310	EST	5.4
KC_181310 M97936	M97936	SIGNAL TRANSDUCER AND ACTIVATOR OF	4.5
DC A A 242757	AA242757	ESTs	4.0
WQ2001	W92001		4.0
RC AA398721	AA398721	EST - RC_AA398721	4.0
DC_A A448410	AA448410	ESIS	4.0
DC_A449419	AA479348	H.sapiens mRNA for SYT	5.4
NC_747473746	C02170	ESTs Weakly similar to weak similarity to ribosomal	5.4
C02110 C02110	A A 437225	ESTs	5.4
NC A45/22	N51917	ESTs	5.4
PC 4 4 293773	AA293773	Homo sapiens clone 23870 mRNA sequence	4.0
RC_A A 449357	AA449357	ESIS	4.0
DC DA1294 c	R41294	ESIS	4.0
pc W45275 f	W45275	CD44 antigen (cell adhesion molecule)	4.0
RC_AA447213	AA447213	ESTs Weakly similar to 50S RIBOSOMAL PROTEIN L20	
RC_AA135809	AA135809	\$183 \$193	, k
RC_AA191524	AA191524	5153 STST	5,5
RC_AA399477	AA399477	PKTS	5.3
RC_H80749	H80749	Tomas camiline olicomeric matrix protein (COMP)	5.3
RC_N94385_s	N94585	FeT - RC A A 157811	5.3
RC_AA157811	AA157811	101 TWO.	

5.3	, v	, ,	, v.	53	5.3	5.3	5.3	53	5.3	5.2	5.2	5.2	5.2	5.2	5.2	5.2	5.2	5.2	5.2	5.2	7.0	7:C	7.0	5.2	5.5	5.2	5.2	5.1	5.1	5.1	5.1	5.1	5.1	5.1	5.1	5.1	5.1	
ESTs	5153 5 4:4 50 5000;if = 5 1;	Stoudie Kindse to Stoudie Kindse to Stoudie St	Homo sapiens Jakz Kinase incara comprete cus	NUCLEAR FORE COMPLETA INCIPATION 21	Homo sapiens gammaz-adapun (Ozazo) muzaka compress	Human Ko/SSA noonacteoptotean monday (1997)	SIST SIST TEST TO STREET TO SERVE TO SE	ESIS WEAKLY SHILLIAI TO DANGE LED TO SEE	Human low-Mr G.F-buiding protein (1904) 1 transporter of the KINASE	EKBB-2 KECEFION FNOIEM-11100M12 EXPE	CICE (170k) II sluha (170k)	ESTS	Homo saniens DNA sequence from PAC 262D12 on	Homo sapiens U-snRNP-associated cyclophilin (USA-	EST - RC_AA488280	ESTs	EST - RC_N73861	G1 to S phase transition 1	Protein kinase C substrate 80K-H	ESTs	ESIS	ESI	ESIS	ESTs Moderately similar to ZINC FINGER PROTEIN 7	ESI RCT	STOT TOTA	TET TET TO THE TOTAL OF THE TOTAL TET TO	ESAS faightly shinital to 1111 O 1111 City and complete ode			TSH	ESTS		2 A A 4 A 4 A 4 A 4 A 4 A 4 A 4 A 4 A 4		HSTS H	STSH STSH	
AA425154	AA284565	AA287022	AA464860	AA401428	AA394071	AA195036	AA465191	AA476293	US9877	X03363	AA621714	AA026682	AA0053570	A A 403116	A A 488280	H96392	N73861	X17644	H87319	AA452857	R92205	N54321	AA279160	AA599140	AA609891	R05312	R59183	W45302	W59961	AA481453	K42030	AA434152	W60180	N79612	AA487449	R43543	AA284518	N98461
RC_AA425154	RC_AA284565	RC_AA287022	RC_AA464860	RC_AA401428	RC_AA394071	RC_AA195036	RC_AA465191	RC_AA476293	US9877	X03363	RC_AA621714	RC_AA026682	RC_AA025370	RC_AA005262	DC_A4483110	00799407 OU	RC_N73861	100CM ON	H87319	RC AA452857	RC R92205	RC_N54321	RC AA279160	RC_AA599140	RC_AA609891	RC_R05312_s	RC_R59183_f	RC_W45302	RC_W59961_s	RC_AA481453	* RC_R42036	RC_AA434152	RC_W60180	RC_N79612	RC_AA487449	RC_R43543	RC_AA284518	RC_N98461

Y09912 mal	Y09912	H.sapiens mRNA for AP-2 beta transcription factor	5.1
AA491465	AA491465	ESTs	5.1
RC_AA436673	AA436673	ESTs	5.1
N73808 f	N73808	55153 1551	0.0
_ J05614	J05614	ESI - J05614	0.0
RC F10496 f	F10496	H.sapiens 40 kDa protein kinase related to rat ERK2	0.0
_ J003 14	J00314	Homo sapiens clone 24703 beta-tubulin mRNA complete	0.0
RC AA085676	AA085676	ESTs Weakly similar to TYL [H.sapiens]	0.0
_AA236356	AA236356	ESTS	5.0
RC_AA252395	AA252395	ESTs	5.0
C N33927 s	N33927	"Homo sapiens mRNA for histone H2B clone pJG4-5-	0.0
RC AA262179	AA262179	ESIS	0.0
RC_AA211400	AA211400	ESIS	2.0
M80244	M80244	INTEGRAL MEMBRANE PROTEIN E16	5.0
M16336	M16336	CD2 antigen (T cell surface antigen T11)	5.0
AA479995	AA479995	Homo sapiens mRNA for KIAA0583 protein partial cds	0.0
D82419	D82419	ESTs Highly similar to UBIQUITIN-CONJUGATING	0.0
RC AA055892	AA055892	ESIS	0.0
RC_AA281451	AA281451	ESIS	5.0
_ U62392	U62392	Homo sapiens zinc finger protein mRNA complete cds	5.0 5.0
RC AA172056	AA172056	ESIS	0.0
RC_AA287095	AA287095	EST - RC_AA28/095	0.0
RC_AA425691	AA425691	ESIS TOT	0.4
RC_AA426376	AA426376	ES18 HST	0.0
RC_AA446000	AA446000	E013	0.0
RC_AA478951	AA478951	ESI	0.0
RC_F02080_f	F02080	STOTE THE THE STORY OF THE ALTERNATIVE TO THE STORY OF TH	5.0
RC_T79815	C186/.I.	ESIS WEAKIJ SIIIIIIai to :::: CALO SOLI IAIATA I	4.9
RC_F10945	F10945	Folypynmique tract binding protein (macer 1) Ueme equipus clone 23915 mRNA sequence	6.4
C01169	C01109	ESTS	4.9
	AA490830	SESE SESE	4.9
RC_AA133/56	AA135/30	Destain absendatase 2 (formerly 2A) catalytic subunit	4.9
RC_H24044	H24044	FIGURIA PHOSPHARASC 2 (1911) 21/1 Carian processing FIGURIA	4 9
RC_R41772	K41//2	FST - RC T59338	4.9
KC_159338	93536	Cadherin 11 (OR-cadherin)	4.9
D21255	D21233	ESTS	4.9
RC_AA234559	AA234339 M34803	ESTS Highly similar to HYPOTHETICAL 47.8 KD	4.9
KC_N34893	A A 101512	ESTS	4.8
KC_AA191512	21C1CIAA 21C0003W	Human mRNA for KIAA0203 gene complete cds	4.8
, w 0000 v _	0000		

4.8	4.8	4.8	8.4	4.7	4.7	4.7	4.7		. .	. t	4.7	4.6	4.6	4.6	4.6	4.6	4.6	4.6	4.6	4.5	4.5	4.5	4.5	4.5	4.5	4.5	4.5	4.4	4.4	4.4	4.4	4.4	4.4	4.4	4.4	4.4	4. 4	4. 4	4 4.
ESTs	EST	FSTS	TSH	FSTS	FINONI IOIBLE HUAN of actions of the street mon	ESIS Hignly Similal to Indul. Colored Property PSTs	STATE	5. T. C. T.	ESIS	ESTs Weakly similar to CALMODULIN [D.melanogaster]	STORE STORE	HATS.	DAP-3 mBNA	Terrange T 8004 Totaline to 1 8004 Total	EST- RC AA177051	ESTS	A DVI AMINE N. A CETVI TRANSFERASE	ANTERIMENTAL PROPERTY IN THE TRANSPORT OF THE PROPERTY OF THE	ESTS	Total Wookly similar to T0169 4 [C elegans]	ESTS WEARLY SILLING TO LOT CO. T. EST. F.ST.	ESTS	ESTS	EST	Human stromelysin-3 mRNA	Transfer of the solution of th	Homo sapiens spinitic port board protein specific contraction transfer and transfer	FST - RC 197307	Urokinase-type plasminogen activator	ESTs	Human fibroblast activation protein mRNA complete cds	H.sapiens mRNA homologous to S. cerevisiae RAD54	ESTs	ESTs Moderately similar to !!!! ALU SUBFAMILY J	EST	ESTS	ESIS	Homo sapiens mRNA for JM27 protein complete CUS	ESTs Highly similar to CHOLINE DEHYDROGENASE
A A 400513	C1C00+AA	7074207	AA406081	AA448158	AA404352	AA075599	AA026356	AA157836	AA196549	AA417321	AA418074	N32919	AA620795	U18321	H9/017	AA1//031	AA453483	XI/059	M34438	17689N	AA464853	AA210722	AA461507	140841	0/01/N	00//CX	AA255605	AA443634	19/30/	A A 430124	A A 405569	A A 2 2 7 9 0 0	A A 422025	AA346495	AA386260	AA398155	AA453466	AA463726	C20981
	RC_AA400513	RC_N94362	RC_AA406081	RC_AA448158	RC_AA404352	AA075599	RC_AA026356	RC_AA157836	RC_AA196549	RC_AA417321	RC_AA418074	RC_N32919	RC_AA620795	U18321	RC_H97012	RC_AA17/051	RC_AA453483	X17059	M34458_mal	RC_N68921	RC_AA464853	RC_AA210722	RC_AA461507	RC_T40841	RC_N71076	X57766	RC_AA255605	RC_AA443634	RC_T97307	X02419_mai	RC_AA430124	RC_AA403309	KC_AA221300	PC A 346495	BC_AA386260	RC_AA398155	RC_AA453466	RC_AA463726	RC_C20981

4.4	4.4	4. 4 4. 4	£.4	5.4 5.4	J. c	£.3	4. ε	£.4	4.2 2.4	4.2	4.2	4.2	4.2	4.2 2.4	7. C	4. 4. 4. c.	4.7	4.2	4.2	4.1	4.7		4. 4	1.,	4. 4	 -	4.1	4.0	4.0	4.0	4.0	4.0	4.0	4.0	3.0	6.6	3.9	3.0	,
EST	RAG (recombination activating gene) cohort 1	PROTEASOME COMPONENT C9	Human DNA polymerase delta small subunit mRNA	Human protein kinase ATR mRNA complete cds	ESIS	Human cyclin-selective ubiquitin carrier protein mRNA	Human lysyl oxidase-like protein mRNA complete cds	Human mRNA for reticulocalbin complete cds	ESTs	ESTs	ESTs Highly similar to CYTOCHROME P450 IVB1	Cellular retinoic acid-binding protein [human skin mRNA	ESTs	Human cysteine protease Mch2 isoform alpha (Mch2)	EDIS	Human putative M phase phosphoprotein 2 (MPP2) mKNA	EDIS	High-mobility group (nonhistone chromosomal) protein 2	Proliferating cell nuclear antigen	Human mRNA for KIAA0255 gene complete cds	ESIS	SIST COLOR	H.sapiens mRNA for BS69 protein	EDIS	Protein kinase interferon-inducible double stranded RNA	SICH CONTRACTOR OF THE CONTRAC	Annexin XI (30kU autoantigen)		nom, tilakin dimilar to DBDY protein [H sar		Small induvible cytokine A 5 (RANTES)	Siliali lituaciolo ej tormo iso esta esta esta esta esta esta esta esta	Cytochrome B561		TT TO THE TOTAL STATE OF THE TOT	Homo sapiens nuclya 101 A 104 Protection Protection PSTP	A DD wikewlation fac	•	ENTS Weakly similar to the delinition thre round
B70801	1728386	AA206497	U21090	AA453176	AA235112	U73379	U24389	D42073	H18947	H90161	H25577	S74445	AA419200	AA227959	AA416931	U74612	R46482	X62534	M15796	D54296	AA398369	AA464707	AA478799	AA496369.	U50648	N66818	AA448347	AA193297	AA28/390	AA228026	AA421041	75//8W	X94563	U29463	AA287325	R44709	AA256837	W455/2	AA482224
י 1000בת הת	KC_K/0801_S 1128386	RC AA206497	171090	RC AA453176	RC_AA235112	1173379	1124389	D42073	RC H18947	PC H90161 s	RC H25577	S74445	RC AA419200	RC_AA227959	RC_AA416931	_ U74612	RC R46482	X62534	M15796	RC D54296 f	RC AA398369	RC_AA464707	RC_AA478799	RC_AA496369	U50648	RC N66818	RC_AA448347	AA193297	RC_AA287596	RC_AA228026	RC_AA421041	RC_W87752_s	X94563_xpt2_r	U29463	RC_AA287325	RC_R44709	RC_AA256837	RC_W45572_f	RC_AA482224

0, 0, 6, 6 0, 0, 8, 6	o ∞ ∞ ∞ ∞	3.8	. w	 	0 K	3.8	3.8	ي د د د	3.7	3.7	3.7	3.7	3.7	3.7	3.7	3.7	3.6	3.0	3.0	3.6	3.6	3.6	3.6	3.0 2.6	0.0	3.5	3.5	3.5	3.5
ESTs EST ESTs Weakly similar to D9481.16 gene product	Homo sapiens mRNA for SPOP	ESTs Weakly similar to nemo form II [D.melanogaster]	Human mRNA for prepro cortistatin like peptide complete Homo sapiens Amplified in Breast Cancer (AIB1) mRNA	Homo sapiens tumorous imaginal discs protein Tid56	ESTS FSTS	EST	ESTs Moderately similar to The KIAA0138 gene product	Human mRNA for KIAA0078 gene complete cds	ESTS Weakly similar to !!!! ALU SUBFAMILL I FSTs	Human 60-kdal ribonucleoprotein (Ro) mRNA complete	ESTS	ESTs Highly similar to HETEROGENEOUS NUCLEAR	ESTS HQTs	Homo sapiens importin-alpha homolog (SRP1gamma)	Decorin	ESTs	Homo sapiens mRNA for high mobility group protein	ESTs Moderately similar to unknown protein [H.sapiens]	Homo sapiens mKNA for NB thymosin beta complete cus	ESIS Weakly similar to :::: ALU SUBI AMALI ST ESIS	ESTs	ESTs Weakly similar to hypothetical L1 protein	ESTs	ESTS	1531 	Homo sapiens KB07 protein mKNA partial cds	CALCAL CALCALLA TANAMATAN	ESTS Weakly Similar to Lpaop [Secretarians]	H.sapiens mRNA for synaptonemal complex lateral
AA416627 AA485360 AA459960	AA224324 AA132366	AA399244 AA465094	N41018 W46488	N98525	N74501	AA423632 AA485451	AA133527	AA287642	AA232183	N6/102	AA424486	W45728	AA399547	AA598661 V12394	M14219	H04339	AA435840	F02450	N91887	AA401758	N63873	N67603	AA461492	AA521240	N93967	D82558	N36835	AA159181	AA398450
RC_AA416627 RC_AA485360 RC_AA459960	RC_AA224324 RC_AA132366	RC_AA599244 RC_AA465094	RC_N41018 W46488	RC N98525	RC_N74501	RC_AA425652	RC_AA483431 RC_AA133527	RC_AA287642	RC_AA232183	RC_N67102_s	M23077 RC A A 474486	RC_W45728	RC_AA399547	RC_AA598661	I 12394 M14219	RC H04339	RC AA435840	RC_F02450	RC_N91887_s	RC_AA401758	RC_W/3/88	RC_N67603	RC AA461492	RC_AA521240	RC N93967	_D82558	RC_N36835	RC_AA159181	RC_AA126951 RC_AA398450

RC AA112063	AA112063	ESTs Weakly similar to PRE-MRNA SPLICING	3.5
- R70167	R70167	SINI	5.5
RC AA034365	AA034365	NUCLEAR PORE GLYCOPROTEIN P62	3.5 5.6
_ X75346	X75346	H.sapiens mRNA for MAP kinase activated protein kinase	5.5 7.5
3 AA083069	AA083069	EST - RC_AA083069	5.5 7.6
ZAA404593	AA404593	ESIS	5.5
A A 412739	AA412739	EST	3.5
A A 447626	AA447626	EST	3.5
RC_AA453787	AA453787	Human TFIIB related factor hBRF (HBRF) mRNA	3.5
AA599106	AA599106	EST - RC_AA599106	3.5
C H72283 s	H72283	Human mRNA for KIAA0265 gene partial cds	3.5
RC N90859	65806N	ESTS	3.5
_L38961	L38961	Integral transmembrane protein 1	3.5
X69141	X69141	FARNESYL-DIPHOSPHAIE	ر. د. ر
RC R63734	R63734	ESIS	5.0
AA164209	AA164209	Homo sapiens RRM RNA binding protein Gry-rbp (Grk Y-	4. 6
RC_AA449417	AA449417	Homo sapiens mRNA for putative glucosyltransterase	4. c
RC H88639	H88639	ESIS	4.0
RC_AA411448	AA411448	ESIS	4. c
AA258203	AA258203	ESIS	4. c
RC_AA100470	AA100470	ESIS	ų, c
RC N29740	N29740	ESIS	4.0
RC_N34895	N34895	ESTS	4.6
RC AA442070	AA442070	Phosphoribosyl pyrophosphate amidotransferase	κ. 4. 4
_AA115397	AA115397	Homo sapiens mRNA for putative methyltransterase	4.5
RC F10326 f	F10326	EST	3.4
RC N33920	N33920	H.sapiens mRNA for diubiquitin	κ. 4. 4
RC AA429917	AA429917	ESTS	4.4
_AA453164	AA453164	EST	4.0
RC_AA029042	AA029042	Human hSIAH2 mRNA complete cds	ئ 4. د
_ L47276	L47276	EST - L4/2/6	4.6
007806	O07806	DNA topoisomerase I	3.4
S81003	S81003	T-UBC	3.4
RC AA232535	AA232535	ESTs Weakly similar to LINE-1 REVERSE	w (
RC_AA490899	AA490899	ESTs	3.3
RC R45356	R45356	Homo sapiens cDNA similar to RNA binding protein C.	
RC AA047896	AA047896	ESIS	າ ເ ດີ ເ
RC_F09353	F09353	Homo sapiens sodium/myo-inositol cotransporter	
RC_N67437	N67437	LOI S	
RC_AA116036	AA116036	5213	

	, e,	
Human kinesin-like spindle protein HKSP (HKSP) mRNA ESTs ESTs Weakly similar to ZK1058.4 [C.elegans] ESTs Homo sapiens DNA from chromosome 19p13.2 cosmids ESTs EST - RC AA435847 EST - RC N34059 ESTs Weakly similar to !!!! ALU SUBFAMILY SC ESTs Weakly similar to F28F8.3 [C.elegans] ESTs Weakly similar to HYPOTHETICAL 139.1 KD ESTs	Homo sapiens mRNA for KIAA0583 protein partial cds ESTs Homo sapiens polyadenylate binding protein-interacting Peroxisomal membrane protein 3 (35kD Zellweger ESTs Peroxisomal biogenesis factor 7 Human WS-3 mRNA complete cds Human mRNA for KIAA0312 gene partial cds ESTs ESTs ESTs Weakly similar to Pin1 protein [H.sapiens] ESTs	HETEROCHROMATIN PROTEIN 1 HOMOLOG ESTS ESTS Weakly similar to RAR-RESPONSIVE PROTEIN Human siah binding protein 1 (SiahBP1) mRNA partial cds Human IAP homolog B (MIHB) mRNA complete cds Homo sapiens signal recognition particle 72 (SRP72) ESTs EST - RC_N78572 Human myogenic repressor I-mf (MDFI) mRNA complete ESTs ESTs ESTs ESTs
AA453159 AA496051 W85861 R24237 AD000092 W44735 AA435847 N34059 N58172 AA421213 N35385 T15665 W46255	AA490969 F09315 AA169379 AA211941 M86852 AA134965 R60192 D84145 Z97054 AA232939 T65797 W33134 AA609423	L07515 R67996 AA42079 U31586 U37547 U81554 T40327 AA448213 AA047036 W72967 N69331 W44928
RC_AA453159 RC_AA496051 RC_AA496051 RC_R24237_f AD000092_cds RC_W44735 RC_M34059 RC_N34059 RC_NS8172 AA421213 RC_N3585 RC_N3585 RC_N35858 RC_N35858	RC_AA490969	RC_R67996 RC_R67996 RC_R67996 U31586 U31547 U81554 T40327_s RC_N78572 RC_AA448213 RC_AA448213 RC_AA6047036 RC_W72967 RC_W44928 RC_AA150043

3.1.	3.1	3.1	3.1	3.1	3.1	3.1	3.1	3.0	3.0	3.0	3.0	3.0	3.0	3.0	3.0	3.0	3.0	3.0	3.0	3.0	3.0	2.9	2.9	2.9	2.9	6.2	6.7	6.7	2.9	2.9	V. C.	6.7 6.0	6.7	2.9 2.9).7 0 0	7.7
ESTs Homo sapiens ribonuclease P protein subunit p20 (RPP20)	ESIS	Homo sapiens clone 25392 mikha sequence	NOCEEOSIDE AND Anatahyroid adenomatosis 1)	Cyclin Di (1101) parami rece e e e e e e e e e e e e e e e e e	ESTs	Human mRNA for KIAA0276 gene partial cds	EST	PTB-ASSOCIATED SPLICING FACTOR	ESTS	HKR-11	H.sapiens mRNA for TAFII100 protein	Homo sapiens mRNA for nucleolar protein nNop50 FSTs	ECT. Westly similar to CSOF4 12 IC elegans	ESTS Weakly summer to cook in the ESTS	ESTs	ESTS	ESTs Weakly similar to ZK1058.5 [C.elegans]	ESTs Highly similar to ribosome-binding protein p34	H.sapiens Cetg mRNA for chaperonin	Interferon (gamma)-induced cell line protein 10 from	ESTs	Homo sapiens M962 protein spliced isoform 2 mRNA	ESTs	ESTs	Homo sapiens mammaglobin B precursor mRNA complete	ESIS	ESIS	ESIS	ESTs Weakly similar to coded for by C. elegans cDNA	ESTs Weakly similar to W02D9.2 [C.elegans]	Human mRNA for KIAA0252 gene partial cds	EST - RC_AA426120	DNA-BINDING PROTEIN NEFA PRECURSOR	Human tyrosyl-tRNA synthetase mRNA complete cds	elota elotan self. I amo to to	H.sapiens mRNA for ras-related GTP-binding protein
D60208 AA401687	N21626	S66431	X17620	84/62X	AA2328/4	AA291239 D87466	A A 398360	X70944	AA099719	S50223	AA251738	Y12065	AA4005//	N4/204	W80482	AA423027	AA421934 W46286	A A 386764	X74801	AA152305	R36548	N32333	N36881	H16790	AA393164	AA399164	AA164293	D60061	AA203523	AA195936	T25732	AA426120	AA485214	U40714	AA213506	R50840
RC_D60208_f	RC N21626	S66431	X17620	X59798	RC_AA232874	RC_AA291259	D8/400	V70044 s	RC AA099719	\$50223	RC AA251738		RC_AA406577	RC_N47204	RC_W80482		RC_AA22/932	KC_W40200_S	X74801	A/4801 PC AA152305	0075147 ON	RC_N32333	PC_N36881	PC_H16790	A A 393164 s	RC AA399164	RC_AA164293	PC D60061 s	A A 203523	RC AA195936		RC AA426120	RC_AA485214	U40714	RC AA213506	RC_R50840

2, 2, 2, 2, 5, 5, 5, 5, 5, 5, 5, 5, 5, 5, 5, 5, 5,	2.9 2.9	2.9	2.9	2.9	2.9	2.9	0 7 0 8	2.8	2.8	7.0 7.0 7.0	2.8	2.8	7.8 7.8	2.8	2.8	2.8 8.0	6 % 0 %	2.8	2.8	2.8	2.8	2.8 8.0	6.0 0.0	, c % %	2.8	
ESTs Homo sapiens GTP-binding protein (RAB4) mRNA ESTs Highly similar to deduced protein product shows Homo sapiens splicing factor (CC1.3) mRNA complete cds ESTs	ESIS ESTS RSTs	ESTS ESTS FSTs	ESTS ESTS FST - HG110-HT110	Thymidylate synthase	ESTs Weakly similar to 108A11.2 [C.elegans] ESTs	ESTs Moderately similar to rabkinesin-6 [M.musculus]	ESTs Weakly similar to ORF YORZ85W [S.cerevisiae]	H. Sapiens mixed 101 M-phase prosprior mega HEAT SHOCK 70 KD PROTEIN 1	ESTs	TST TST	ESTs Weakly similar to !!!! ALU SUBFAMILY J	Homo sapiens eps8 binding protein e3B1 mRNA complete	ESTS Westing to Similar to Scenerisiae	ES IS Weakly summan to comment to ESTS	EST - RC_T63857	ESTS	Natural resistance-associated macrophage protein 2	ESTS	Human mRNA for clathrin-like protein complete cds	EST - RC_AA446572	ESTs	ESTs Weakly similar to elastin like protein	Uracil-DNA glycosylase	PUTATIVE 60S RIBOSOMAL PROTEIN	Human neutral amino acid transporter Dinguya Complete Human splicing factor SRp30c mRNA complete eds	
	AA242834 F13779	AA206088 AA292747	AA400/25 H97677	TIGR - HG110- D00596	AA279420	AA179845	AA411532	AA292/65 AA609501	AA478596	N48715	AA412497 AA480103	AF006516	AA128407	AA425606 AA232231	T63857	AA488432	R49327	AA405512	A A 310779	AA446572	AA486407	R39234	AA425900	F03605	U53347 U30825	
RC_R97040 M28211 AA452011 RC_AA228020 T54762_s	RC_AA242834 RC_F13779	RC_AA206088 RC_AA292747	RC_AA400725 RC_H97677_s	HG110-HT110 D00596	RC_AA279420	RC_AA179845 RC_AA179845	RC_AA411532	RC_AA292765 BC_AA600501	RC_AA478596	RC_N48715	RC_AA412497 BC_AA480103	_AF006516	RC_AA128407	RC_AA425606 BC_AA23231	RC T63857	RC AA488432	RC_R49327	RC_AA405512	KC_K/961/	RC_A446572	RC_AA486407	RC R39234 r	RC AA425900	RC_F03605_f	U53347 1130825)

RC_N21159	N21159	Homo sapiens forkhead protein (FKHRL1) mRNA	2.8
RC_H24460_s	H24460	(WACC) + minor of the continuing process of the continuing of the continuing process of the cont	, c ×
RC_D80710_f	D80710 X76732	ESIS Weakly similar to transmemorate protein (1.1.34ptens) DNA-BINDING PROTEIN NEFA PRECURSOR	2.8
AA001409	AA001409	ESTS	2.8
RC N92915	N92915	ESTS	2.8
RC_N29325	N29325	ESTs Highly similar to 47 KD PROTEIN. [Pseudomonas	2.8
AA609200	AA609200	EST - RC_AA609200	0.7
$\overline{R}C_{L}R41933$	R41933	HSTS:	2.0 2.0
AA262768	AA262768	ESIS	7.0 0.0 0.0 0.0
RC_R46025	R46025	SISE TSE	0.7
AA164687	AA164687	ES 13	2.7
AA424031	AA424031 N72663	ESTS	2.7
W60160	W69160	ESTs	2.7
RC_W09100 BC_N05798	N25798	ESTs	2.7
AA431333	AA431333	ESTs Highly similar to Ras inhibitor [H.sapiens]	2.7
RC R54112	R54112	ESTs	2.7
RC_F02863	F02863	ESTs Moderately similar to !!!! ALU SUBFAMILY SQ	2.7
W80750	W80750	EVIS	7.7
AA461509	AA461509	ESTs Weakly similar to putative p150 [H.sapiens]	7.7
AA620586	AA620586	EDIS	7.7
L11669	L11669	Human tetracycline transporter-like protein indaya	, , ,
AA291269	AA291269	EDIS FINE	2.7
RC_W87747	W8//4/	T cell recentor alpha-chain	2.7
K02777	K02777	ICIN S A INC	2.7
98668X	9868X	H.sapiens mKNA for NBK apoptous muusei protein	7.7
AA125969	AA125969	ESTS Weakly Similar to FSSOTA: For the participation of the participatio	7.7
AA449718	AA449/18 D02354	ES IS Weakly similar to zing finder, independent to ESTs	2.7
KC_K02534	PCCZONI PCCZONI	ESTs Weakly similar to serine protein kinase SRPK1	2.7
AA423/23	R71481	ESTs	2.7
D63391	D63391	Human mRNA for platelet activating factor	2.7
AA126743	AA126743	ESTs	2.7
U33052	U33052	Human lipid-activated protein kinase PRK2 mRNA	2.7
M96982	M96982	SPLICING FACTOR U2AF 35 KD SUBUNII	7.7
X16396	X16396	NAD-DEFENDENT	7.7
L12350	L12350	7 Infombospondin 2 FCTs	2.7
AA215333	AA215333		27
_AA102520	AA102520	ESTS Highly similar to HYPOTHETICAL 31.9 AND	i

2.56 2.66 2.66 2.66 2.66 2.66 2.66 2.66	5 5 5 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6	, , , , , , , , , , , , , , , , , , ,	; ; ; ; ; ; ; ; ; ; ; ; ; ; ; ; ; ; ;	2 2 2 2 2 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6	2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2
ESTs ESTs Weakly similar to PROCOLLAGEN ALPHA 2(IV) ESTs Homo sapiens 15 kDa selenoprotein mRNA complete cds ESTs Highly similar to elastin like protein	Radin blood group Human mRNA for KIAA0175 gene complete cds Homo sapiens (clone CÇ6) NADH-ubiquinone EST - RC AA412112	Human terminal transferase mRNA complete cds ESTs Human IEF SSP 9502 mRNA complete cds ESTs Homo sapiens IPL (IPL) mRNA complete cds H.sapiens mRNA for spermine synthase	Human mRNA for transcriptional activator hSNF2b ESTs ESTs Weakly similar to semaphorin C [M.musculus] ESTs Homo sapiens mRNA expressed in osteoblast complete cds Human mRNA for KLAA0160 gene partial cds ESTs Weakly similar to ZK1058.4 [C.elegans] ESTs	Homo sapiens nibrin (NBS) mRNA complete eds ESTs Moderately similar to !!!! ALU SUBFAMILY SX Homo sapiens nibrin (NBS) mRNA complete eds Human breast cancer estrogen regulated LIV-1 protein ESTs ESTs	ESTS ESTS ESTS Matrix Gla protein ESTS ESTS ESTS ESTS EST - RC_D60374_f EST - RC_AA069547
N68869 W85712 AA176121 AA001402 N51316	L03411 D79997 L04490 AA412112 AA417956	AA453624 N63210 N92948 AA447553 AF001294 Z49099	AA598648 AA447617 AA293300 AS0333 AB000115 D63881 T39176 N46252	N33516 N48790 H98655 AA242758 AA130349 AA262491 N70646	AA490882 AA610073 N67187 AA404957 T70541 AA236489 AA284372 D60374 AA069547
RC_N68869 RC_W85712 RC_AA176121 RC_AA001402 RC_N51316	L03411 D79997 L04490 RC_AA412112 RC_AA417956	RC_AA453624 RC_N63210 RC_N92948_s RC_AA447553 AF001294 Z49099	RC_AA598648 RC_AA447617 RC_AA293300 RC_RS0333_i AB000115 D63881 T39176_s RC_N46252	RC_N33516 RC_N48790 RC_H98655 RC_A242758 RC_AA130349 RC_AA262491 RC_AA262491	RC_AA490882 RC_AA610073 RC_N67187_s RC_AA404957 RC_T70541 RC_AA236489 RC_AA284372 RC_AA284372 RC_AA069547

2.5	7,4	2.7	C.7	C.2	2.5	2.5	5.5 5.0	2.2 5.0	2.5			2.5	2.5	2.5	2.5	2.5			2.5					2.5	2.4				2.4	2.4	2.4	2.4			2.4	2.4	4.7 4.0	† •
ESTs	SICO.	EST	EDIS	ESIS	KERATIN TYPE II CYTOSKELETAL 6D	STORE SPORE	E) CJ	1631 time of the formal formation of the	rotein phosphatase 4 (rothicity A) catalytic succint	Human micha tol alabout o gene company to gene continua	FSTS	CDC28 protein kinase 1	Human mRNA for KIAA0097 gene complete cds	GTPase-activating protein ras p21 (RASA)	H.sapiens mRNA for Sm protein G	LARGE FIBROBLAST PROTEOGLYCAN	ESTs	ESTS	Homo sapiens mRNA for KIAA0530 protein partial cds	Human mRNA for KIAA0389 gene complete cds	ESTS	ESTS	ESTs	VITAMIN K-DEPENDENT GAMMA-CARBOXYLASE	Homo sapiens diphthamide biosynthesis protein-2 (DPH2)	ESTs Highly similar to ribosome-binding protein p34	ESTS	ESTS	ESTS	EST - RC_AA258601	Human mRNA for KIAA0240 gene partial cds	U2 SMALL NUCLEAR RIBONUCLEOPROTEIN A'	ESTs Highly similar to CHROMOSOME	ESTS	ESI	EST - RC_AA487207	ESTs Weakly similar to F08G12.1 [C.elegans]	פיוטת
AA398280	N93000	AA291503	AA446100	W47183	L00205	AA369027	N66158	AA428179	X/0218	AB002308	AA4168//	X54941	14364G	M23379	X85373	U16306	T39763	H12634	AA251587	AA160890	N21677	AA191424	AA451707	AA045083	AA252672	0906LM	AA262651	AA399047	AA456646	AA258601	N51260	X13482	AA504223	R37778	W31919	AA487207	AA599674	240898
RC_AA398280	RC_N93000	RC_AA291503	RC_AA446100	RC W47183	_ L00205	RC_AA369027	RC_N66158	RC_AA428179	X/0218	AB002308	RC_AA416877	KC_AA202/30 V5/041	774741	M23379	X85373	1116306	T39763 s	RC H12634	RC AA251587	RC_AA160890	RC N21677	RC AA191424	RC_AA451707	RC_AA045083	RC_AA252672	0906LM	RC AA262651	RC_AA399047	RC_AA456646	RC_AA258601	RC N51260 s	X13482	AA504223	RC R37778	RC W31919	RC AA487207	RC_AA599674	RC_Z40898

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2, 2, 2, 2, 4, 4, 4, 4, 4, 4, 4, 4, 4, 4, 4, 4, 4,	2.2.2.2.4.4.4.4.4.4.4.4.4.4.4.4.4.4.4.4	1	12.2.2.2.2.4.4.4.4.4.4.4.4.4.4.4.4.4.4.4	2 2 2 2 2 2 4 4 4 4 6 6	5 5 3 3 3 3 3 3 3 5 5 5 5 5 5 5 5 5 5 5	23 23 23 23 23 23 23 23 23 23 23 23 23 2
EST - RC_AA286942 Human Rho-associated coiled-coil containing protein ESTs Weakly similar to D9481.16 gene product ESTs Weakly similar to U1 SMALL NUCLEAR ESTs	Human mRNA for KIAA0079 gene complete cds HLA-DR ASSOCIATED PROTEIN I COL10A1	Guanylate binding protein 1 interferon-inducible 67kD H.sapiens mRNA for UDP-GalNAc:polypeptide N-Glutamine-fructose-6-phosphate transaminase Human mRNA for KIAA0242 gene partial cds	Human mKNA for 3-announdazote-1-carboxannuc-1- ESTS Homo sapiens mRNA for ATP-dependent RNA helicase ESTS Human serine kinase mRNA complete cds	ESTS ESTS Highly similar to VACUOLAR ATP SYNTHASE ESTS ESTS Highly similar to CHROMOSOME ESTS Moderately similar to ZINC FINGER PROTEIN	Human protein-tyrosine phosphatase (HU-PP-1) mRNA ESTs Homo sapiens breast cancer putative transcription factor ESTs Weakly similar to R01H10.8 [C.elegans] CDC28 protein kinase 2 ESTs Highly similar to HYPOTHETICAL TRP-ASP ESTs	ESTs Weakly similar to PROBABLE UBIQUITIN ESTs ESTs Highly similar to GAG POLYPROTEIN [Avian ESTs ESTs Moderately similar to !!!! ALU SUBFAMILY SQ ESTs Human mRNA for EBI1-ligand chemokine complete cds
AA286942 AA371604 AA223209 AA218663	D38555 D38555 U73477 X60382	X84373 M55542 S82597 M90516 D87684	D82348 AA190993 N69352 AA434329 T29681	AA279799 T25896 AA147708 AA039887 AA455239 Z39909	AA330771 AA173223 R81830 AA010065 W23625 F02907 N04581	AA069285 AA450116 N33063 T87807 W49574 AA250737 AA425749 U77180
RC_AA286942 RC_AA371604 RC_AA23209 RC_AA218663	XC_AA449458 D38555 U73477 X60382_ma1	X84373 M55542 S82597_ma1 M90516 D87684	D82348 RC_AA190993 RC_N69352 AA434329 RC_T29681_f	RC_AA279799 RC_T25896 RC_AA147708 RC_AA039887 RC_AA455239 RC_AA455239	AA330771_s RC_AA17323 RC_AA01814 RC_AA010065 RC_W23625_s RC_W23625_s RC_W23625_s	RC_AA069285 RC_AA450116 RC_N33063 RC_T87807_s RC_W49574 RC_AA425749 RC_AA425749

C AA024658	AA024658	ESTs	2.3
_ D00591	D00591	Chromosome condensation 1	C.7
X94453	X94453	Pyrroline-5-carboxylate synthetase (glutamate gamma-	2.3
RC_AA459673	AA459673	ESTs Highly similar to CHROMOSOME	2.3
.C_AA428647	AA428647	83052 7g F35	23
RC_R52088	R52088		23
X54199	X54199	Phosphoribosylglycinamide tolinylitatistetase	2.3
RC_N49284_s	N49284	MITS FROID-ONO-CANA COMPANY	23
U37022_ma1	U37022	Human cyclin-dependent protein kinase mouth compared	2.3
D26156	D26156	Human mKINA Iof transcriptional acuvator maint 20) c
U72514	U72514	Human Czi inkiya compicie cus	, c
S79873	S79873	Lysosomal-associated memorane protein 2	., c
U47077	U47077	Human DNA-dependent protein kinase catalytic subunit	4.C
U59423	U59423	Human chromosome 4 Mad homolog Smad1 mKNA	2.2 2.3
J05633	105633	Integrin beta-5 subunit	£.2
Z48042	Z48042	H.sapiens mRNA encoding GPI-anchored protein p13/	2.3
A A 037657	AA037657	ENIS	2.3
RC N29888	N29888	Human NAD+-specific isocitrate dehydrogenase beta	2.3
C A 4751776	AA251776	ESTs	2.3
C_A A 282568	AA282568	ESTs Weakly similar to F25H2.6 [C.elegans]	2.3
2C_A A 236951	AA236951	ESTs	2.3
C A A 464423	AA464423	ESTs Weakly similar to !!!! ALU SUBFAMILY J	2.3
C_A A 037410	AA037410	Human DNA sequence from PAC 127B20 on chromosome	2.3
R63652	R63652	ESTs	2.3
DC_N66857	N66857	ESTs	2.3
7500VI VI	A A 280588	ESTs	2.2
XC_AA260366	A A 436477	ESTs	2.2
0.0000 V V	A A 028028	ESTs	2.2
D N20148	N39148	ESTs	2.2
RC_1439140	A A 485773	ESTs	2.2
C_AA403223	A A 011556	ESTs	2.2
C AAUITOO	967630V V	ESTs	2.2
3C_AA053636	AA033030	ESTS	2.2
C_AA148516	AA148510	FCTs	2.2
3C_AA495924	AA495924	ESTS	2.2
C AA131692	AA131692	STORY STORY	2.2
RC N90401	N90401	ES1S TST	2.2
RC AA436613	AA436613	SO 10.2 Committee	7.7
RC_AA397921	AA397921	Homo sapiens mKNA transcriptional unit 143	2:2
RC_AA251766	AA251766	ESTS Moderately similar to metastasis-associated gene	2.2
RC F09328	F09328	LO 10	1

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2	2.2	7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7	2:2	5.2 5.2 5.3 7.7 7.7 7.7	2.2	2.2	2.2	2.2	2.2	2.2	2.2	2.2	2.2	2.2	2.1	2.1	2.1	2.1	
ESTS ESTS ESTS ESTS ESTS	ESTs Human DNA from overlapping chromosome 19 cosmids ESTs	ESTs ESTs Moderately similar to neurexophilin 2 [M.musculus] EST - RC_AA490237	H.sapiens mRNA for HOXC9 protein exon 1 ESTs	Cyclin-dependent kinase inhibitor 2A (melanoma p10 ESTs Weakly similar to ZNF127-Xp [H.sapiens] ESTs	ESTs EST - RC_AA430726	ESTs ESTs	Human cysteine-rich heart protein (hCRHP) mRNA ESTs Weakly similar to Diff33 gene product [H.sapiens]	ESTs Home canions Am2/3 protein complex subunit p20-Arc	Glycyl-tRNA synthetase	Human 26S proteasome-associated pad1 homolog (POH1) COATOMER BETA' SUBUNIT	Homo sapiens protein regulating cytokinesis 1 (PRC1) ESTs	ESTs Weakly similar to ISOLEUCYL-TRNA	ESIS Homo sapiens vesicle transport related protein mRNA	ESTs Highly similar to UBIQUITIN-CONJUGATING ESTs	ESTs ESTs	ESTs Highly similar to YSA1 PROTEIN [Saccharomyces	Homo sapiens RRM RNA binding protein Gry-rbp (Grx r- ESTs Moderately similar to !!!! ALU SUBFAMILY SP	ESTs Human mRNA for kinesin-related protein partial cds	
T15674 AA416735 AA423827	H84658 H89261 H99261 T10060	AA043160 AA059214 AA490237	AA227856 H28428	AA076328 F13690 AA287320	AA287833 AA430726	N93618 T59686	U09770 R72008	N94606	U09510	U86782 X70476	AA417030	AA236516	R78119 AA150088	AA043353 AA126719	AA403121 N64378	AA158132	AA253031 Z99394	AA400820 W20391	
RC_T15674_f RC_AA416735 RC_AA423827	RC_H84658_s RC_H99261_s RC_T10060	AA043160 RC_AA059214 RC_AA490237	RC_AA227856 RC_H28428	RC_AA076328 RC_F13690_s BC_AA287320	RC_AA287833 RC_AA430726	RC_N93618	009770 U09770 RC R72008	RC_N94606	KC_D80237_s U09510	U86782 X70476	RC_AA417030	RC_AA236516		RC_AA043353	RC_AA403121 PC N64378	RC_AA158132	RC_AA253031 RC_799394_s	RC_AA400820 RC_W20391 s	

22.2.2.2.2.2.2.2.2.2.2.2.2.2.2.2.2.2.2.2	22.1	22222	2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7	2 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5
ESTs Weakly similar to HYPOTHETICAL 46.4 KD H. sapiens RNA for CLCN3 Human COP9 homolog (HCOP9) mRNA complete cds Human Hs-cul-4A mRNA partial cds PTB-ASSOCIATED SPLICING FACTOR Human CDP-diacylglycerol synthase (CDS) mRNA ESTs	Homo sapiens MAD3-like protein kinase mRNA complete ESTs ESTs Weakly similar to similar to S. cerevisiae LAG1 ESTs	Cut (Drosophila)-like 1 (CCAAT displacement protein) ESTs Weakly similar to !!!! ALU SUBFAMILY SP ESTs ESTs ESTs ESTs	ESTS ESTS ESTS Human mRNA for KIAA0331 gene complete cds ESTS ESTS Human tubulin-folding cofactor E mRNA complete cds	ATP-DEPENDENT DNA HELICASE II 86 KD Homo sapiens mRNA for GDP dissociation inhibitor beta Human ubiquitin-homology domain protein PIC1 mRNA SRY (sex determining region Y)-box 4 ESTs LARGE FIBROBLAST PROTEOGLYCAN Homo sapiens testis-specific nm23 homolog mRNA EST	Homo sapiens clone 24651 mRNA sequence Homo sapiens vacuolar H(+)-ATPase subunit mRNA ESTs ESTs ESTs Homo sapiens SKB1Hs mRNA complete cds ESTs Highly similar to SIGNAL RECOGNITION
AA281780 R91380 U51205 U58090 X70944 U60808	AAA251909 AA134063 H38246 AA451712	H80/3/ M74099 AA018587 AA291137 AA426660 AA427662 AA45148	AA610039 N72113 W32470 AA620464 AA609869	M30938 D13988 U67122 X70683 N22222 AA393695 W37384	N95837 N95837 N24968 AA598452 AA287388 AA487202 F02651 AF015913 AA476582
RC_AA281780 RC_R91380_s U51205 U58090 X70944 U60808	AA460077 RC_AA251909 RC_AA134063 RC_H38246_s RC_AA451712	RC_H80/3/s M74099 RC_AA018587 RC_AA291137 RC_AA426060 RC_AA427662 PC_AA46148	RC_AA403148 RC_AA610039 RC_N72113 RC_W32470 RC_AA620464 RC_AA609869	M30938 D13988 U67122 X70683 RC_N22222 RC_AA393695	RC_N95837 RC_N95837 RC_N24968 RC_AA598452 RC_AA287388 RC_AA487202 RC_F02651 AF015913 RC_AA476582

2:0 2:0 2:0 3:0	5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5	5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5	2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7	0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0
ESTS ESTs ESTs Weakly similar to PEREGRIN [H.sapiens] ESTs	PHOSPHATIDYLINOSITOL 3-KINASE CATALYTIC ESTS ESTS ESTS ESTS ESTS ESTS	H.sapiens mRNA for novel member of serine-arginine ESTS ESTS ESTS Human mRNA for KIAA0393 gene complete cds Homo sapiens mRNA for osteoblast specific cysteine-rich Homo sapiens retinoblastoma-associated protein HEC EST - RC_H11938	ESTS Highly similar to Zinc Linger protein parameters. ESTS ESTS Highly similar to PRE-MRNA SPLICING FACTOR ESTS CAD PROTEIN Human selenium donor protein (selD) mRNA complete cds ESTS	Homo sapiens unknown mRNA complete cds ESTs Acid phosphatase 1 soluble Human p55CDC mRNA complete cds ESTs ESTs	H.sapiens mRNA for I.BA2 protein Androgen receptor (dihydrotestosterone receptor testicular Aldehyde dehydrogenase 8 EST - D28364 Human mRNA for KIAA0225 gene partial cds H.sapiens polyA site DNA Human hepatocyte nuclear factor-3 alpha (HNF-3 alpha)
R68425 W80467 D53392 Z39053	N91246 Z29090 AA443460 AA045481 W28362 AA233177	H78241 AA443596 AA453255 X69636 AA047265 AA188981 A11938	123539 AA405838 AA426375 N24954 D78586 U34044 AA621122 Z40810	AA127716 AA237022 AA479139 U05340 AA417909 AA181657 L37347	X92896 M23263 U37519 D28364 D86978 Z24724 U39840
RC_R68425 RC_W80467 RC_D53392_f RC_Z39053	RC_N91246 	RC_H78241_s RC_AA443596 RC_AA453255 RC_AA453255 RC_AA047265 RC_AA188981 RC_AA188981	RC_T23539 RC_AA405838 RC_AA426375 FC_N24954 D78586 U34044 RC_AA621122 PC_Z40810	RC_Z40810 RC_AA127716 RC_AA237022 RC_AA479139 RC_AA417909 RC_AA1181657 L37347	X92896 M23263 U37519 D28364 D86978 Z24724 U39840

5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5	2.0 2.2.0 1.2.0 9.9 9.9	v e e e e e e e e e e	9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9	61 61 61 61 61 61 61 61
ESTs ESTs ESTs Human mRNA for KIAA0178 gene partial cds Homo sapiens meltrin-L precursor (ADAM12) mRNA ESTs	ESTs Core-binding factor beta subunit H.sapiens mRNA for RNA polymerase II subunit Human mRNA for clathrin coat assembly protein-like	ESTs Highly similar to FK506-BINDING PROTEIN ESTs H.sapiens mRNA for transcriptional intermediary factor 2 ESTs Weakly similar to No definition line found Homo sapiens thyroid hormone receptor-associated protein Human mRNA for KIAA,0020 gene complete cds	ESTs Highly similar to HYPOTHETICAL 16.5 KD ESTs ESTs ESTs ESTs ESTs ESTs ESTs ESTs	ESTs Weakly similar to siah binding protein 1 [H.sapiens] H.sapiens mRNA for 2.19 gene ESTs Highly similar to UBIQUITIN-CONJUGATING ESTs Weakly similar to T13F2.1 [C.elegans] ESTs Weakly similar to Y53C12A.3 [C.elegans] ESTs Highly similar to HYPOTHETICAL 37.2 KD Signal recognition particle 19 kD protein Homo sapiens ICT1 (alias DS-1) mRNA Murine leukemia viral (bmi-1) oncogene homolog Human protein kinase mRNA complete cds
AA136884 T63174 N50963 D80000 R73567 N70520	AA476312 AA521474 L20298 Z47727 U91932 AA173417	AA28/854 W80763 AA112679 AA233261 T77464 AA227463 D50920 D13645	AA233168 AA227963 AA451898 AA30745 N23393 W37933 AA504832 AA446918	T89703 X55448 AA257972 Z40715 AA464013 W95063 X12791 X81788 L13689 L13689
RC_AA136884 T63174_s RC_NS0963 D80000 RC_R73567 RC_N70520	RC_AA476312 RC_AA521474 L20298 U91932 RC_AA173417	RC_AA287834 RC_W80763 RC_AA112679 RC_AA233261 RC_T77464 RC_AA227463 D50920	RC_AA233168 RC_AA227963 RC_AA451898 RC_AA302745 RC_N23393 RC_AA504832 RC_AA46918 RC_AA46918	RC_T89703 X55448_cds1 RC_AA257972 RC_Z40715 AA464013 RC_W95063 X12791 X81788 L13689 L33801

1.9	6.1	1.9	1.9	1.9	1.9	1.9	V.1.	1.9	1.9	1.9	1.9	1.9	1.9	6: F	P.1.	1.9	∞ ∞	8	1.8	1.8	1.8	»: «	× :	8. 0	o «	× ×	8	1.8	1.8	1.8	8: 6	0.1	1.0
Human BRCA1-associated RING domain protein EST - HG174-HT174	Human fetal Alz-50-reactive clone 1 (FAC1)	Small nuclear ribonucleoprotein polypeptide B	ESTS	ESTs	EST	Human heterochromatin protein HP1Hs-gamma mRNA	Human Gu protein mRNA partial cds	ESIS	ESTS FSTs	CATHEPSIN K PRECURSOR	ESTs	PLASMA-CELL MEMBRANE GLYCOPROTEIN PC-1	ESTS	ESTs	ESTS	ESTS	ESTs Moderately similar to !!!! ALU SUBFAMILY SQ	II cms sminn mDMA for n115 complete cds	HOING Sapicins that A to a forth to the South to the Sout	ESTs Weakly similar to F25D7.1 [C.elegans]	Homo sapiens clone 23770 mRNA sequence	ESTS	ESTs Moderately similar to ALR [H.sapiens]	Human mRNA for KIAA0208 gene complete cds	ESTS RCT	NIPTO GO SEI ELICA GO GO CALLERON CONTRACTOR	ESIS Weakly similar to FROBABLE ESIGNOTEIN	ES18 Modelately summat to oos read coordinate from ESTs	ESTs	ESTs Highly similar to DOSAGE COMPENSATION	Centromere protein A (17kD)	ESIS	518
U76638 TIGR - HG174-	X58072 U05237	M21259	AA149585	A A 236453	T25867	U26312	U41387	AA496000	AA489046	AAZ/0023 V82153	AA403008	D12485	H44386	N46423	N55336	AA497052	N26855	N52006	240332	AA279171	AA251982	AA455001	AA599219	W84790	H94248	AA234765	N35583	AA099241	A A 420088	F02990	U14518	AA232103	AA398319
U76638 HG174-HT174	X58072 U05237	M21259	RC_AA149585	AA115058_S	KC_AA230433	U26312	U41387	RC_AA496000	RC_AA489046	RC_AA2/8653	A04133		H44386 s	RC N46423	RC_N55336	RC AA497052	RC_N26855	RC_N52006	RC_Z40332	RC_AA028074		AA455001 s	RC AA599219	RC_W84790_s	RC_H94248	RC_AA234765	RC_N35583		KC_AA430192	KC_AA420988	U14518	RC AA232103	RC_AA398319

1.8	 8	1.8	1.8	%: %:	8. .	. .	8.1	1.8	1.8	1.8	1.8	1.8	1.8	1.8	1.8	1.8	1.8	1.8	8:1.8	7.8	8.I. 8	3.5	×	×. 1.	0.1	0.0	0.1	0.1	× -	0.0	0.1	0.1	v. i	0.0	1.0
ESTs Gastrin-releasing peptide	Homo sapiens RRM RNA binding protein Gry-rbp (GRY-	EST - RC_T96595	Homo sapiens mRNA for Dnm1p/Vps1p-like protein	Homo sapiens Werner syndrome gene complete cds	Exostoses (multiple) 2	Human protein phosphatase (KAPI) mKNA complete cds	FSTS	ESTS	ESTs	OX40L RECEPTOR PRECURSOR	ESTs Highly similar to transcription factor ARF6 chain B	ESTs Highly similar to GERANYLGERANYL	ESTs	ESTs Moderately similar to HYPOTHETICAL 52.2 KD	Human RNA-binding protein CUG-BP/hNab50 (NAB50)	Human transducin beta-2 subunit mRNA complete cds	H.sapiens mRNA for transmembrane protein mp24	Human transportin (TRN) mRNA complete cds	Threonyl-tRNA synthetase	Damage-specific DNA binding protein 1 (127 kD)	V-jun avian sarcoma virus 17 oncogene homolog	Homo sapiens splicing factor (CC1.3) mRNA complete cds	Homo sapiens DNJ3/CPR3 mRNA complete cds	ESIS	ENIS	ESTS Weakly similar to COLH6./ [C.elegans]	ESIS	SICT	ESIS	Homo sapiens spleen mitotic checkpoint BUB3 (BUB3)	ESIS	ESISE CONTRACTOR OF THE PROPERTY OF THE PROPER	Human mRNA for KIAA0005 gene complete cds	HETEROGENOUS NUCLEAR RIBONUCLEUPROTEIN	ESTs Weakly similar to SOF1 PROTEIN [Saccharomyces
W52225 AA063460	AA44/603 AA401274	F04989 T96595	AF000430	L76937	U72263	L25876	W68502	W72876	AA457566	X75962	AA232104	AA417962	D59894	AA598988	W88983	M36429	X92098	U70322	M63180	U32986	U65928	L10910	R41281	N50050	AA460350	AA292066	AA291923	AA411144	AA436171	AA251758	AA406478	AA191353	D13630	X65488	AA131584
RC_W52225 RC_AA063460	RC_AA447603 RC_AA401274	RC_F04989 RC_T96595	AF000430	L76937_ma1	U72263	L25876	W68502	EC AA005108	RC AA457566	X75962	RC AA232104	RC_AA417962	RC D59894	RC AA598988	RC W88983	_M36429	X92098	U70322	M63180	U32986	U65928	L10910	RC R41281	RC_N50050	RC_AA460350	RC_AA292066	RC_AA291923	RC_AA411144	RC_AA436171	RC_AA251758	RC_AA406478	RC_AA191353	D13630	X65488	RC_AA131584

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ESTs Moderately similar to YY1-associated factor 2 ESTs Weakly similar to W04D2.6 [C.elegans] ESTs ESTs Human GABA-A receptor pi subunit mRNA complete cds ESTs ESTs ESTs ESTs Homo sapiens clone 23797 and 23917 mRNA partial cds ESTs Human arginine-rich nuclear protein rec.22b mRNA ESTs Human arginine-rich nuclear protein mRNA complete cds ESTs Human arginine-rich nuclear protein mRNA complete cds ESTs Human arginine-rich nuclear protein mRNA complete cds ESTs	Ħ
AA283743 AA056588 AA180321 AA262957 AA262957 AA234767 AA470961 T03865 N51226 AA470140 AA459005 AA470140 AA459005 AA470140 AA458900 N31598 T57317 W55890 N31598 T57317 W55890 N31598 AA111879 AA111879 AA111879 AA111879 AA410972 W23469 AA282315 R39923 AA465690 AA465690 AA465690 AA465690	L19101 U90551 L76703 TIGR - HG4557-
RC_AA283743 RC_AA056588 RC_AA180321 RC_AA262957 RC_AA479961 RC_AA479961 RC_AA479961 RC_AA479961 RC_AA479961 RC_AA47629 RC_AA47629 RC_AA470140 RC_AA470140 RC_AA470140 RC_AA425439 RC_AA416075 RC_AA111879 RC_AA116075 RC_AA116077	L19161 U90551 L76703 HG4557-

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Cathepsin B ESTs Chaperonin containing T-complex subunit 6 ESTs ESTs ESTs ESTs ESTs ESTs Proto-oncogene AML1 {alternative products} ESTs ESTs ESTs ESTs ESTs ESTs ESTs ESTs	Homo sapiens mRNA for KIAA0286 gene partial cds EST ESTS Highly similar to KINESIN-LIKE PROTEIN KIF4 ESTS H.sapiens mRNA for unknown protein expressed in ESTS ESTS Weakly similar to L8004.7 gene product Homo sapiens cyclin T2a mRNA complete cds Homo sapiens mRNA for GDP dissociation inhibitor beta Human enhancer of zeste homolog 2 (EZH2) mRNA EST ESTS
N58561 H15436 L27706 W85888 AA479362 D31161 AA035143 AA416733 N9222 AA416733 N9222 AA410894 AA609053 AA400080 AA400080 AA135814 AA46522 AA415803 AA135814 AA45222 AA45222 AA43052 AA456437	AA424524 AA252360 W58247 AA599622 X89059 W04698 N73865 T23820 AA099589 U61145 AA151708
RC_N58561_s RC_H15436 L27706 RC_MA479362 D31161_s RC_AA4035143 AF002668 RC_AA416733 RC_N92860_s RC_AA416733 RC_AA416733 RC_AA416894 RC_AA469053 RC_AA400080 RC_AA400080 RC_AA460053 RC_AA41773 RC_AA41773 RC_AA465222 RC_AA465437 s	RC_AA24524 RC_AA252360 W58247_s RC_AA599622 X89059 RC_W04698 RC_N73865 RC_T23820 RC_AA099589 RC_AA151708 RC_AA151708

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9.1	1.6	1.6	1.6	1.6	1.6	9.1	1.6	1.6	1.6	1.6	1.6	1.6	1.6	1.6	1.6	1.6	1.6	1.6	1.6	1.6	1.6	1.6	1.6	1.6	1.6	1.6	1.6	1.6	1.6	1.6	1.6	1.6	1.6	1.6	1.6	1.6	1.6	9.7	1.6
ESTs	EST	ESTs	Glycerol-3-phosphate dehydrogenase 2 (mitochondrial)	ESTs	ESTs	ESTs Weakly similar to Yel007c-ap [S.cerevisiae]	ESTs	Homo sapiens SOX22 protein (SOX22) mRNA complete		Human mRNA for KIAA0345 gene complet	ESTs	Homo sapiens DNA polymerase zeta catalytic subunit	Human mRNA for KIAA0110 gene complete cds	Homo sapiens interferon stimulated T-cell alpha	PROBABLE 26S PROTEASE SUBUNIT TBP-1	ESTs	ESTs Weakly similar to NADH-UBIQUINONE	Human mRNA for ceramide glucosyltransferase complete	Nuclear autoantigenic sperm protein (histone-binding)	Human nuclear factor NF45 mRNA complete cds	Tumor protein p53 (Li-Fraumeni syndrome)	Human non-histone chromosomal protein (NHC) mRNA	NAD(P)H:menadione oxidoreductase	MULTIFUNCTIONAL PROTEIN ADE2	Human Tat-SF1 mRNA complete cds	X-LINKED HELICASE II	NUCLEAR PORE GLYCOPROTEIN P62	ESTs Weakly similar to ZK1058.4 [C.elegans]	ESTs	ESTs	Human clone 23722 mRNA sequence	ESTs	ESTs Highly similar to hypothetical protein 100K	ESTs	Homo sapiens mRNA for KIAA0530 protein partial cds	Cathepsin B	ESTS	ESTs	Homo sapiens UDP-glucose dehydrogenase (UGDH)
AA417970	R10720	AA256106	U12424	W73820	AA167708	AA187579	R15743	N69014	Z38919	AB002343	AA521186	AA258205	D14811	U59286	M34079	AA026418	N26259	D50840	M97856	U10323	M22898	U90549	103934	X53793	U76992	U09820	X58521	AA121127	AA243007	AA463195	00606N	W15528	R70621	AA456598	AA167375	AA279667	F03738	AA148885	D60856
RC AA417970	RC R10720	AA256106	U12424 s	RC W73820	RC AA167708	AA187579	RC R15743	RC N69014 s	RC Z38919	AB002343	RC AA521186	RC_AA258205	D14811	US9286	M34079	RC_AA026418	RC N26259	_D50840	M97856	U10323	M22898	U90549	103934	X53793	U76992	U09820	X58521	RC_AA121127	RC_AA243007	RC AA463195	00000	RC W15528	_R70621	RC AA456598	RC_AA167375	RC_AA279667	RC_F03738_f	RC AA148885	RC D60856 f

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1.6	1.6	1.6	1.6	1.6	1.6	1.6	1.5	1.5	1.5	1.5	1.5	1.5	1.5	1.5	1.5	1.5	1.5	1.5	1.5	1.5	1.5	1.5	1.5	1.5	1.5	1.5	1.5	1.5	1.5	1.5	1.5	1.5	5.5	1.5	1.5	1.5	1.5	1.5	1.5
EST	ESTs	Human fetal Alz-50-reactive clone 1 (FAC1) mRNA	ESTs Highly similar to POP2 PROTEIN [Saccharomyces	H.sapiens p63 mRNA for transmembrane protein	Homo sapiens KIAA0442 mRNA partial cds	Homo sapiens importin beta subunit mRNA complete cds	ESTs	ESTs Weakly similar to DFS70 [H.sapiens]	ESTs	ESTs	ESTs	ESTs	ESTs	ESTs	Protein tyrosine phosphatase non-receptor type 4	ESTs	EST	ESTs Weakly similar to similar to mouse MMR1	ESTs	ESTs	Homo sapiens putative transcriptional repressor E2F-6	ESTs	ESTs	ESTs Highly similar to CALCIUM-TRANSPORTING	Human mRNA for phosphatidylinositol-glycan-class C	Deoxyguanosine kinase	EST	EST - AA113913	ESTs	EST	ESTs	Human mRNA for KIAA0064 gene complete cds	ESTs Highly similar to COATOMER ZETA SUBUNIT	Homo sapiens clone 24800 mRNA sequence	ESTS	ESTS	ESTs Highly similar to CLATHRIN COAT ASSEMBLY	Homo sapiens LIM protein mRNA complete cds	Transcription factor 3 (E2A immunoglobulin enhancer
AA101811	AA347967	U05237	AA256678	X69910	H95039	AA181580	AA219699	AA236672	AA476319	AA041551	AA195179	AA256492	W93640	R07016	AA232644	N37065	T10258	AA279757	N67390	AA489086	W72138	N68640	D51177	AA400271	D85418	U41668	AA133309	AA113913	N21978	AA447970	AA433925	D31764	AA146888	W28366	AA293568	N48677	AA226922	N52271	M31523
RC AA101811	RC_AA347967	U05237	RC AA256678	01669X	RC H95039	RC AA181580	RC_AA219699	RC_AA236672	RC_AA476319	RC_AA041551	AA195179 s	RC AA256492	RC W93640	RC_R07016	RC AA232644	RC N37065	RC_T10258	RC AA279757	RC N67390	RC AA489086	RC W72138	RC_N68640	RC_D51177	RC AA400271	D85418	U41668	RC AA133309		RC N21978	RC AA447970		D31764	AA146888 s	W28366	RC AA293568	RC N48677	RC AA226922	RC N52271	M31523

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1.5	1.5	 	5.	1.5	1.5	1.5	1.5	1.5	1.5	1.5	1.5	1.5	1.5	1.5	1.5	1.5	1.5	1.5	1.5	1.5	1.5	1.4	1.4	1.4	1.4	1.4	1.4	1.4	1.4	1.4	1.4	1.4	1.4	1.4	4	1.4	1.4
Human mRNA for KIAA0077 gene partial cds V-akt murine thymoma viral oncogene homolog 1	EST - HG884-HT884	Human amyloid precursor protein-binding protein i	Homo sapiens neterochroniaum protein p23 mayes	nullan Erd NOT 213-21 markin compress cas	Eukaryotic translation initiation factor 4C (eIF-4C)	ESTs Weakly similar to DIPEPTIDYL PEPTIDASE IV	ESTs	ESTs Weakly similar to HYPOTHETICAL PROTEIN	H.sapiens mRNA for centrin gene	Human serine kinase mRNA complete cds	ESTS	ESTs	ESTs	H.sapiens mRNA for translin	EST - RC_R87660	ESTs Moderately similar to !!!! ALU SUBFAMILY J	Homo sapiens GDP-L-fucose pyrophosphorylase (GFPP)	ESTs Weakly similar to ORF2 consensus sequence	ESTs Weakly similar to DPY-30 PROTEIN [C.elegans]	ESTS	ESTs Highly similar to HYPOTHETICAL 66.5 KD	H.sapiens NAP (nucleosome assembly protein) mRNA	ESTs	ESTs	ESTs Weakly similar to similar to tyrosyl-tRNA	ESTs	Homo sapiens brain expressed ring finger protein mRNA	ESTs Moderately similar to M-phase phosphoprotein 11	ESTs	ESTs	ESTs	Homo sapiens TFAR19 mRNA complete cds	Zinc finger protein 7 (KOX 4 clone HF.16)	ESTS	ESTS	ESTs Highly similar to GTP-BINDING PROTEIN SARA	ESIS
D38521 M63167	TIGR - HG884-	U50939	U35451	U94830 I 33881	L18960	AA496257	AA262942	AA056735	AA482014	U09564	AA609738	N54450	Z39255	N78717	R87660	T98843	Z39211	AA412528	N26101	AA399550	AA621580	AA422160	AA156542	H59417	AA460246	F04982	AA285277	R09196	F09983	AA057193	AA449068	AA452724	M29580	AA417895	AA425100	AA287879	T17440
D38521 M63167	HG884-HT884	U50939	U35451	U94836 1 33881	1.18960	RC AA496257	RC_AA262942	RC_AA056735	RC_AA482014	_ U09564	RC AA609738	RC N54450 i	RC_Z39255_f	RC_N78717_s	RC R87660	RC_T98843	RC_Z39211	RC AA412528	RC_N26101	RC AA399550	RC_AA621580	_AA422160	RC AA156542	H59417 s	RC AA460246	RC F04982	AA285277	RC R09196	RC_F09983	RC AA057193	RC_AA449068	_AA452724	M29580	RC AA417895	RC_AA425100	RC_AA287879	RC_T17440_f

4.1	t. T	1.4	1.4	1.4	4	1. 4.	1.4	1.4	1.4	4	t. T	1.4	1.4	1.4	1.4	1.4	1.4	4	† · ·	1. 4.	1.4	1.4	1.4	4.1	4. 7	1.4	1.4	1.4	1.4	1.4	† ·	1.4 4.1	1.4	
ESTs	ESIST TOTAL	ESTS	ESTs	Cyclin-dependent kinase inhibitor 2A (melanoma p16	Human PMS5 mRNA (yeast mismatch repair gene PMS1	Transcription factor 6-like 1 (mitochondrial transcription FSTs	ESTS	ESTs Weakly similar to ASPARTYL-TRNA	ESTS Weakly similar to !!!! ALU SUBFAMILY SB1	ESTS	ESIS	I dietora rof VMam Classes.		ESTS	H.sapiens HUNKI mRNA	Himan himor necrosis factor type I receptor associated	fidulian tunnor meeting fraction of ESTs	Homo sapiens Sox-like transcriptional factor mRNA	SISE	Homo sapiens chromosome 19 cosmid R32469 FSTs	Tr mp NA for PKII-alpha partial cds	11			ESTs Weakly s			Criston singer determined	DNA mismatch repair protein texts.	ESTs Weakly similar to HYPOTHETICAL 48.		Homo sapiens clone 23965 mRNA sequence		
A A 2 5 5 5 5 4	A A 402937	N92293	AA292128	K01243	026/2/	M62810	AA279991	N66569	AA28/138 A A 278755	AA278733 AA195515	R63925	N23972	W93379	L06419	AA411882	AA083918	012595	AA262945 AA135095	T16226	AA497015	AA171939	AB004884	U84720	240041	738501	H93708	AA025086	N64244	U07418	R64660	0 A 490949	D80921	N26722	W90146
62332CA 4 Oct	KC_AA23337	RC N92293	RC AA292128	RC_R01243	U26727	D38498_1 M62810	RC_AA279991	RC_N66569	RC_AA287138	RC_AA278755	EC AA195515 50 P63975	RC_N23972	RC W93379_s	_ L06419	RC_AA411882	RC_AA085918	U12595	RC_AA262943	KC_AA133093	RC AA497015	RC_AA171939	_AB004884	U84720	RC_Z40041	RC_W60473	KC_229301 pc_Hq3708_s	RC A A 0 2 5 0 8 6	RC N64244			RC_AA400093	RC_AA490949 RC_D80921_s	RC N26722	RC_W90146_f

1.4	. . .	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	E: -	1.3
Homo sapiens mRNA for KIAA0512 protein complete cds	STORE STORE	DI-N-ACETYLCHITOBIASE PRECURSOR	Human protein-tyrosine phosphatase (HU-PP-1) mRNA	ESTs	ESTs	ESTs Weakly similar to C01A2.4 [C.elegans]	ESTs	Homo sapiens HP protein (HP) mRNA complete cds	Homo sapiens mRNA for dachshund protein	ESTs	Homo sapiens clone 23592 mRNA sequence	ESTs	H.sapiens RBQ-1 mRNA	Zinc finger protein 139 (clone pHZ-37)	ESTs	ESTs	Homo sapiens putative transcriptional repressor E2F-6	Human kruppel-related zinc finger protein (ZNF184)	CD47 antigen (Rh-related antigen integrin-associated	ESTs	ESTs	ESTs	ESTs Weakly similar to ZINC FINGER PROTEIN 135	EST	ESTs	ESTs	ESTs	ESTs	ESTs	ESTs	ESTs	ESTs	ESTs	ESTs	ESTs	ESTs	ESTs	ESTs
AA114250	AA031357 768510	M95767	W67524	AA465093	H26417	T64438	D11718	Z41963	AA252079	AA262889	AA487492	AA464428	H38828	H71863	AA040696	AA258189	AA443294	U66561	X69398	AA122394	T40707	AA435536	AA206800	AA001386	AA428992	R49886	T95591	AA338760	AA098864	AA234817	AA059051	AA046619	AA461169	AA449071	N24732	AA400195	AA029264	W86978
RC_AA114250	RC_AA031357	168310 M95767	RC W67524	RC AA465093	RC H26417	RC_T64438	RC_D11718	RC Z41963 r	RC AA252079	RC_AA262889	RC_AA487492	RC_AA464428	RC H38828 s	RC_H71863_s	RC AA040696	RC_AA258189	RC_AA443294	_ U66561	X69398	RC AA122394	RC T40707	RC AA435536	RC_AA206800	RC_AA001386	RC_AA428992	RC R49886	RC_T95591	RC AA338760	RC_AA098864	_AA234817	RC AA059051	RC_AA046619	RC_AA461169	RC_AA449071	RC N24732	RC AA400195	RC_AA029264	RC_W86978

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1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.3	1.2	1.2	7:7	7.1	7:1
Human N-ethylmaleimide-sensitive factor mRNA partial	ESTS	Homo sapiens chromosome 19 cosmid R32469	ESTs Highly similar to Polio virus receptor protein	ESTs	Homo sapiens agrin precursor mRNA partial cds	ESTs Moderately similar to !!!! ALU SUBFAMILY SB	ESTs	Protein phosphatase 2 (formerly 2A) regulatory subunit B	Human putative RNA binding protein RNPL mRNA	Human protein tyrosine kinase mRNA complete cds	EST - U96113	H.sapiens mRNA for TIM17 preprotein translocase	SRY (sex-determining region Y)-box 9 (campomelic	ESTs Moderately similar to !!!! ALU SUBFAMILY SB	ESTs	Homo sapiens RRM RNA binding protein Gry-rbp (GRY-	Down-regulator of transcription 1 TBP-binding (negative	ESTs	ESTs	Human brain secretory protein hSec10p (HSEC10) mRNA	ESTs Highly similar to probable chloride channel 3	ESTs	Homo sapiens mRNA for DDS1beta protein complete cds	ESTS	ESTs	ESTs	Human mRNA for KIAA0276 gene partial cds	ESTs	Human mRNA for KIAA0179 gene partial cds	ESTS	H.sapiens mRNA for translin associated protein X	ESTs Highly similar to RSP5 PROTEIN [Saccharomyces	ESTs Moderately similar to NAD(+) ADP-	Homo sapiens mRNA for KIAA0539 protein complete cds	Integral transmembrane protein 1	ESIS	ESIS	ESTS	ESTS
H38086	F13663	AA458542	AA101601	AA598675	AA156670	AA059415	AA485424	M64929	U28686	U02680	U96113	X97544	Z46629	H60061	AA007234	AA329211	AA132007	AA417569	AA236200	AA126426	AA504499	AA521471	H83438	T23932	T59859	N20630	D87466	AA083339	AA598506	AA278650	AA599718	AA398243	N51855	W52065	L38961	N55304	AA147543	T32794	U51698
RC_H38086	RC_F13663	AA458542	RC AA101601	RC_AA598675	AA156670 s	AA059415	RC AA485424	_ M64929	U28686	U02680	U96113	X97544	Z46629	RC_H60061	RC_AA007234	AA329211_s	RC_AA132007	RC_AA417569	RC_AA236200	RC_AA126426	RC_AA504499	RC_AA521471	RC_H83438_s	RC_T23932_f	RC T59859	RC N20630 i	D87466	AA083339	RC_AA598506	RC_AA278650	RC_AA599718	RC_AA398243	RC_N51855	RC_W52065_f	L38961	RC_NS5304_s	AA147543	RC_T32794_s	U51698

ESTS Neuroblastoma RAS viral (v-ras) oncogene homolog ESTS Homo sapiens clone 23685 mRNA sequence CCAAT/enhancer binding protein (C/EBP) gamma ESTS ESTS	
AA464758 X02751 W87544 AA025746 R62444 H05635 AA433943 N42440 W03007 T99364 AA411708 U20240 AA112222 W61011 AA010188 N67104 N71027 AA398222 T85190 N74635 Z38839 U79718 AA355201 Z14077 N68622	T17498
RC_AA464758	RC_T17498

Ratio tumor v. breast	133.3	10/.3	81.8	75.3	72.9	61.5	57.1	53.9	53.2	52	46.8	45.7	38.9	33.0	0.70	52.4	31.1	30.6	28.3	28.1	78	25.5 23.9	22.8	21.7	20.9	19.5	19.4	17.1	17.1	16.8	16.8	16.2	16.1	16.1	
	ESTS	_		EST - RC AA461510	Collagen type XI alpha 1	ESTs	EST - RC N27351	TI TOTAL TOTAL STATE OF STATE OF THE PROPERTY	H.saplens in A. S. C.	focal adhesion kulase (1772)	ESTs	ESTs	ESTs	ADVI AMINE N-ACETYLTRANSFERASE	EST - RC_AA232294	EST - RC_R86839	EST	EST	ESTs	EST - RC_AA211831	ESTs	Inhihin heta A (activin A activin AB alpha polypeptide)	EST RSTs	1212 1212 1212 1212 1212 1212 1212 121	EST - RC_AA232345 ESTs	ESTs	EST. Moderately similar to POL POLYPROTEIN [Feline	ESIS	CARCINOEMBRYONIC ANTIGEN PRECURSOR	ESTS	TOT DC AA211158	EST	ESTs	ESTs Highly similar to thyroid disease hypothetical	
Gene Name								**		Human focal adhesion Ku	ESIS Weakly sum			TWA TVOA								Inhihin heta A (activi					nore Moderately simil	ES IS INOCCIACO	CARCINOEMBE	<u>.</u>					
	Accession	T79956	AA453638	AA461322	AA461510	R67275	AA453518	N27351			AA285050	AA291468	Z40805	AA169440	D90041	AA232294	R86839	AA453641	AA609955	AA283905	AA211831	AA412090	621/2N 621/39	R03/02	K9/003	AA463189	AA421171	AA2518/5	AA054228	AA621402 A A 505133	A A 488191			AA196768 H83527	
	A ffsmetrix ID	DC T79956	50011 N	RC_AA453656	RC AA461510	KC AR401313	KC_K0/2/5	KC AA+33310	KC_N2/331	pC_AA453479	RC_A A 285050	pc_AA291468	DC 740805	RC AA169440	D90041 s		RC AA23223	NO 4 4 453641	KC_AA403041	PC_A 4783905	DC_A A 211831	RC_AA412090	RC_N27159_s	RC_R65763	RC_R97063	RC_AA232940	RC_AA421171	RC_AA251875	RC_AA054228	RC_AA621462	RC_AA505155	RC_AA488191	RC_AA211158	RC_AA196768 H83577 s	110001

16.1	15.8	15.8	15.2	14.9	14.7	14.6	14.6	14.4	14.4	14.4	14.3	14.2	14.1	41	14	13.9	13.9	13.7	13.5	13.4	13.3	13.3	13.3	13.1	13.1	13	12.8	12.5	12.5	12.4	12.4	12.3	12.3	12.2	12.2	12.1	12.1
EST - RC_AA196721	Homo sapiens clone 23967 unknown mRNA partial cds	Inhibin beta A (activin A activin AB alpha polypeptide)	ESTs	ESTs Weakly similar to B0334.4 [C.elegans]	Human fibroblast activation protein mRNA complete cds	ESTs	ESTs	ESTs Moderately similar to PTTG gene product	ESTs	Carboxypeptidase B1 (tissue)	ESTs Highly similar to HYPOTHETICAL 21.5 KD	ESTs	Chromogranin A (parathyroid secretory protein 1)	ESTs Moderately similar to 25E8.1 [D.melanogaster]	ESTs Weakly similar to TH1 protein [D.melanogaster]	Androgen receptor (dihydrotestosterone receptor testicular	ESTs Weakly similar to hypothetical protein 1 [H.sapiens]	ESTs	Homo sapiens histone macroH2A1.2 mRNA complete cds	ESTs Weakly similar to 52-kD SS-A/Ro autoantigen	Human mRNA for KIAA0007 gene partial cds	Homo sapiens mRNA for SCGF-beta complete cds	ESTs	ESTs Moderately similar to !!!! ALU SUBFAMILY SC	EST - RC_AA412065	Homo sapiens regulator of G-protein signalling 12	ESTs	Homo sapiens mRNA for A+U-rich element RNA binding	ESTs	EST	ESTs	ESTs Weakly similar to !!!! ALU SUBFAMILY SX	Homo sapiens ES/130 mRNA complete cds	ESTs	ESTS	ESTS	Fibroblast growth factor receptor 2 (bacteria-expressed
AA196721	T25875	XS7579	AA191404	AA262969	AA436611	R51309	AA461297	AA430032	AA280679	M81057	R07976	R46627	AA461559	AA092129	AA436893	M23263	AA486538	D20379	AA076138	AA045074	D60354	T78922	AA401334	AA431350	AA412065	R61740	R54950	AA037285	AA233796	AA219305	AA252245	AA041276	AA463874	AA461528	AA099404	AA214305	AA220223
RC AA196721	RC T25875	X57579	RC AA191404	RC_AA262969		RC R51309	RC AA461297	RC_AA430032	RC_AA280679	_ M81057	RC R07976	RC_R46627	RC AA461559	AA092129 f	RC AA436893	_ M23263	RC AA486538	RC D20379	RC AA076138	RC_AA045074	RC D60354 s	RC_T78922_s	ĀA401334	RC AA431350	RC_AA412065	RC R61740 f	RC R54950	AA037285	RC AA233796	RC_AA219305	RC_AA252245	RC_AA041276	RC_AA463874	RC_AA461528	RC_AA099404	RC_AA214305	AA220223

e 12.1						_	_	s 11										•	_			A 9.9				se 9.7							ls 9.3			(s 9.2
Glutamine-fructose-6-phosphate transaminase	ESTS	ESTS	UBIQUITIN-LIKE PROTEIN GDX	ESTS	ESTs Weakly similar to extracellular protein [H.sapiens]	Homo sapiens chromosome 9 P1 clone 11659	EST	ESTs	EST	Homo sapiens U4/U6 small nuclear ribonucleoprotein	EST	ESTS	ESTs Weakly similar to MOESIN/EZRIN/RADIXIN	Human splicesomal protein (SAP 61) mRNA complete cds	GANGLIOSIDE GM2 ACTIVATOR PRECURSOR	EST	ESTs Weakly similar to dynein 74K chain cytosolic	ESTs Highly similar to ZINC FINGER PROTEIN 85	ESTS	ESTs Weakly similar to trabecular meshwork inducible	ESTs Weakly similar to No definition line found	H.sapiens DAP-3 mRNA	ESTs	ESTS	ESTS	Spermidine synthase	ESTS	ESTs Weakly similar to R01H10.6 [C.elegans]	PROTEASOME COMPONENT C13 PRECURSOR	EST - RC_T9734	ESTs Highly similar to HYPOTHETICAL 23.1 KD	ESTS	ESTs	ESTS	Human mRNA for KIAA0101 gene complete cds	ESTS
AA478571	AA253217	AA470074	103589	R22952	W56363	AA179298	AA449232	AA281733	AA452601	AA035630	AA453630	R40431	AA405098	AA599259	X62078	AA253170	AA470156	AA281290	AA449832	AA427898	AA609867	R49198	AA112396	AA207015	R06986	M34338	AA228030	AA447982	Z14982	T97341	AA143190	AA282914	N21678	AA262111	D14657	AA007344
RC_AA478571	RC_AA253217	BC A A 236010	103589	RC R22952 s	RC W56363	RC AA179298	RC_AA449232	RC_AA281733	RC_AA452601	RC_AA035630	RC_AA453630	RC R40431	RC AA405098	RC_AA599259	X62078	RC AA253170	RC_AA470156	RC_AA281290	RC_AA449832	RC_AA427898	RC_AA609867	RC R49198 i	RC AA112396	RC_AA207015	1	M34338	_	RC_AA447982	Z14982 mal	RC T97341	RC AA143190			RC AA262111	_ D14657	RC_AA007344

9.2	9.1	6	8.9	∞ t	/.×	×.	0.0	. « . «	2.8	8.4	8.4	8.4	8.4	8.3	8.2		∝	7 0	7.0	7.9	7.9	7.9	7.9	7.9	7.9	7.8	7.8	7.8	×. c	×. c	0.7		1:1
ESTs Highly similar to TRYPSINOGEN ANIONIC ESTs	Human CENP-F kinetochore protein mRNA complete cds	EST - FIGESON -	\$100 calcium-binding protein A7 (psoriasin 1)	H.sapiens Humig mRNA	ESTS	ESIS	ESIS	EDIS	Human mkina ioi niaansi 14 gene panna cus Collagan tune XI alpha 1	Users of the state	Prohibitin	ESTs	EST - RC_T65004	ESTs	Matrix metalloproteinase 2 (gelatinase A collagenase type	STON FOU	ESIS TACH	E2013	SICA CAMPACA C	ESTs Weakly similar to LINE-1 KEVEKSE EST	A / F coli RecA homolog RAD51	Keplication protein A (E con nech mullioned ESTs	ESTs	ESTs	ESTs	EST's Weakly similar to HYPOTHETICAL 88.1 KD	Homo sapiens Ran binding protein 2 (RanBP2alpha)	EST - RC_AA342084	EST	EST - RC_N46435	ESIS	SICH	Homo sapiens mRNA for tyrosyl sulfotransferase-2
W73140	AA447666	TIGR -	K38919 M86757	X72755	AA443342	AA481281	AA608723	AA457018	AA113011	H96237	585655	92666N	T65004	N93197	105070	R40177	D60302	W93659	N69464	AA458882	AA421/50	N33011	PA44/3/4	047748 4 4 7 2 5 0 0 0	P01634	A 150182	A A 446486	AA342084	AA609170	N46435	AA417213	T88814	AA459389
RC_W73140	RC_116508_1 RC_AA447666	HG2981-	RC_K38919_1 M86757	X72755	RC AA443342	RC_AA481281	RC_AA608723	RC_AA457018	RC_AA113011	RC_H96237_s	KC_AA024833	20000X 7 G	DC_T65004	NC_103004	J05070	RC R40177	RC_D60302	RC_W93659	RC_N69464	RC_AA458882	RC_AA421750	RC_N33011_s	RC_AA44/5/4	KC_K4/948_1	RC_AA235009	KC_K01034	RC_AA150182	RC_A A 342084	RC_AA609170	RC N46435	RC AA417213	RC T88814	RC_AA459389

7.6	7.6	7.6	7.6	7.6	7.5	ر: د ر	 	7.5	7.5	4.7	4.7	4.7	7,4	7.4	7.3	7.2	7.2	7.2	7.2	7.1	7.1	7.1	7.1	7.1	7.1	7.1	<i></i>	- 1-	, 69	60	, «	8.9	8 9	8.0	6.7	
ESTs Highly similar to EPIDERMAL GROWTH	ESTs	Human melanoma antigen p15 mRNA complete cds	EST - HG2981-HT3938	EST	Homo sapiens mRNA for osteoblast specific factor 2 (OSF-	Human mRNA for KIAA0170 gene complete cds	ESTs	EST - W01296	ESTS	Interferon (gamma)-induced cell line protein 10 from	EQ 13	ESTS TST	ESTS	FSH	ESTS	FSTS	asenaphyoonom faminosimist and Alka	Homo sapiens mkina iof kynuteimie 5-monosaygemase FST - RC AA427950	ECT. Weath similar to 1111 ALU SUBFAMILY J	ESTS	ESTs Weakly similar to T04A8.11 [C.elegans]	ESTS	H. sapiens mRNA for melanoma growth regulatory protein	ESTS	ESTs	ESTs	ESTs Weakly similar to T06D8.5 [C.elegans]	Š	Homo sapiens clone 23913 indays Homo sapiens clone 23913	STOTE CONTRACTOR OF THE PROPERTY OF THE PROPER	Receptor protein-tyrosine Kinase EDDAN	ESTS	ti niemob obdodoombert mitstern is it is it in men	ESTs Highly similar to purative hydrophobic domain in	numan cione 25509 munan Collagen tyne V alpha	L
H99879	T68871	019796	TIGR -				AA476937	W01296	AA282074	X02530	N67889	AA609309	AA412477	AA459392	AA599042	AA445/94	AA121315	R65593	AA42/950	AAU88438	1401239	A A 236177	A A 282143	AA283003	A A 421158	T10082	Z40345	AA310499	N34686	N71704	U48705	AA419461	AA411204	AA346385	D51229	M11/18
07 H09879	DC_T68871	T119796	HG2981-	3003DIA 74	D13666	RC A A 454566	PC A 4476937	W01296	BC AA282074	X02530	RC N67889	RC_AA609309	RC_AA412477	RC_AA459392	RC_AA599042	RC_AA443794	RC_AA121315	RC_R65593_s	RC_AA427950	RC_AA088458		AA31090/_s	RC_AA230177	RC_AA282143	DC_A A 421158	DO T10082 f	RC Z40345	RC AA310499	RC_N34686	RC N71704	U48705_ma1	RC_AA419461	RC_AA411204	RC_AA346385	RC_D51229_f	M11718

6.7	6.7	6.7	6.7	6.7	6.7	9.0	9.9	6.5	6.5	6.4	6.4	6.4	6.4	6.3	6.3	6.3	6.2	6.2	6.2	6.2	6.2	6.1	6.1	6.1	6.1	6.1	6.1	6.1	6.1	6.1	6.1	6.1	0.1	0 9	0
Homo sapiens mRNA for Efs1 complete cds Human autoantigen pericentriol material 1 (PCM-1)	ESTs	Human bumelanide-sensinye iva-xo conansporez	Human JTV-1 (JTV-1) mRNA complete cds	ESTs Moderately similar to unknown protein [H.sapiens]	Interleukin 6 signal transducer (gp130 oncostatin M	ESTS	ESTS	ESTs Highly similar to COP1 REGULATORY PROTEIN	EST	4	Human pyridoxal kinase mRNA complete	EST	ESTs	H.sapiens mRNA for RNA polymerase II subunit			Human NADH:ubiquinone oxidoreductase subunit B13	ESTs	ESTs	ESTs	Protein-tyros		Human eukaryotic translation initiation factor (eIF3)	Tubulin gamma polypeptide	ESTs	ESTs Highly similar to GOLIATH PROTEIN [Drosophila	ESTs			Human germline oligomeric matrix protein (COMP)	ESIS	ERGIC-53 PROTEIN PRECURSOR	ESTs Weakly similar to line-1 protein ORF2 [H.sapiens]	Collagen type XI alpha 1	ESI
N50550 L27841	T92935	U30246 A A 034069	U24169	AA435849	H99935	R51988	AA505141	AA236384	AA431085	X54326	90968N	AA195651	AA430211	D81608	L77701	AA443658	AA024664	AA279943	AA098874	AA412106	U40271	L37747	U78525	T77733	D20280	W69807	AA133199	H55748	AA479933	L32137	AA448349	AA600257	R99978	J04177	AA406137
RC_N50550 _L27841	RC_T92935	U30246 BC A A 034069	U24169	RC AA435849	RC H99935_s	RC_R51988	RC AA505141	_AA236384	RC AA431085	_ X54326	90968N	RC AA195651	RC_AA430211	D81608	L77701	RC AA443658	RC_AA024664	RC_AA279943	RC_AA098874	RC_AA412106	U40271	L37747 s	U785 <u>2</u> 5	RC T77733_s	RC D20280	RC W69807	RC AA133199	RC H55748	RC AA479933	_ L32137	RC AA448349	RC_AA600257	RC_R99978	J04177	RC_AA406137

9 0 4	5.9	5.8	5.8	5.8	5.8	5.8	5.8	5.8	5.7	5.7	5.7	5.7	5.7	7.0	7.7	5.7	5.0	0.0	0.0	2.4	5.0	5.6	0.0	0.0	2.4	ر. د د	, v	5.5	, v		4.0	4. v	4.0	4.0	4.0	4.0
Homo sapiens short chain L-3-hydroxyacyl-CoA	STSH STSH	Himan extracellular matrix protein 1 (ECM1) mRNA	ESTs	Cyclin B1	ESTs	ESTs	ESTs	ESTs	EST - HG2981-HT3125	Fibronectin 1	ESTs Highly similar to G2/MITOTIC-SPECIFIC	Homo sapiens mRNA for follistain-related protein (FRP)	Homo sapiens mRNA for KIAA0585 protein partial cds	FACTOR VIII INTRON 22 PROTEIN	ESIS	ESIS	ESIS	131 - U9134/	ENIS	CLCA.	ESTs Moderately similar to TYKi protein [M.musculus]	H.sapiens DAF-1 mKINA EST:	STOT	Guanine nucleotide binding protein (G protein) beta	che chellenne direct	Human fibroblast activation protein mkINA complete cus		Homo sapiens Chromosome 10 BAC cloud canada and a sapient control of the same sapients canada and sapients	T2.) Ongoadenyiale synustase are	ESIS	FOL	Laminin receptor (2H5 epitope)	SIGNAL TRANSDUCER AND ACTIVATOR OF	EST - RC_AA398/21	ESIS	ESIS
U73514	W38407	AA292655 1765032	003932	M25753	CC/CZIAI CO/OCC V V	767617W	A A 47 5379	A A 422007	TIGR -	R02572	A A 442763	A A 149624	AA459945	M34677	AA454562	W30943	AA232956	U91327	AA453987	AA040154	T23528	X76105	AA398212	AA416986	T95057	U09278	AA443602	AA075200	X02874	R43883	T81310	H75933	M97936	AA398721	AA448410	AA242757
U73514	RC_W38407	RC_AA292655	756500	KC_AA405/40	CC/C2MI	RC_AA219292	KC_AA28/003	RC_AA423373	RC_AA422007	19670H DC 1903672	27C20X - XI	RC_AA442/03	RC_AA459945	M34677	RC AA454562	W30943	RC AA232956	_ U91327	RC AA453987	RC_AA040154		_X76105	RC AA398212	RC_AA416986	RC T95057 f	_ U09278	RC_AA443602	RC_AA075200	X02874	RC R43883	RC_T81310	RC H75933 i	M97936	RC AA398721	RC_AA448410	RC_AA242757

5.3 5.3	5.3	5.3	5.3 5.3	5.2	5.2	5.1	5.1	1.5	, vo	S	S	5	5	5	5	\$	4.9	4.7	4.7	4.6	4.6	4.5	5.5	4.	4.4	4.4	4.3	4 .3	4.3	4.2	4.2	4.2	4
H.sapiens mRNA for SYT ERBB-2 RECEPTOR PROTEIN-TYROSINE KINASE	Human low-Mr GTP-binding protein (KAB31) mixaya Human germline olisomeric matrix protein (COMP)	Thymidine kinase 1 soluble	Homo sapiens Jak2 kinase mRNA complete cds	ESI - ACAMOND CONTROL OF THE CONTROL	Homo sapiens U-snrur-associated cyclopium (SCTs	Human mRNA for KIAA0389 gene complete cds	EST - RC_AA48/449	5153 515401 T23	FINAL	ESTS	ESTS	ESTs	ESTs	Homo sapiens mRNA for KIAA0583 protein partial cds	ESTs	ESTs	Homo sapiens clone 23915 mRNA sequence	ESTs Highly similar to NADH-UBIQUINONE	ESTS	ARYLAMINE N-ACETYLTRANSFERASE	H.sapiens DAP-3 mRNA	Human stromelysin-3 mRNA	ESTs Weakly similar to T01G9.4 [C.elegans]	H.sapiens mRNA homologous to S. cerevisiae RAD54	ESIS	ESTs Moderately similar to !!!! ALU SUBFAMILY J	Human cyclin-selective ubiquitin carrier protein mRNA	Human lysyl oxidase-like protein mRNA complete cds	Human DNA polymerase delta small subunit mRNA	ESTs Highly similar to CYTOCHROME P450 IVB1	Cellular retinoic acid-binding protein [human skin mRNA	ESTs	ESTs
AA479348 X03363	U59877	N94363 A A 287022	AA464860	AA488280	AA403116 H96392	W59961	AA487449	R43543	J05614	AA2021/9	AA261431	AA423091	A A 446000	A A 470005	A A 055892	AA172056	C01169	AA075599	AA026356	X17059	U18321	X57766	AA464853	AA227900	AA422025	AA346495	173379	1124389	U21090	H25577	S74445	AA419200	AA193297
RC_AA479348 X03363	U59877	RC_N94385_s	RC_AA464860	RC_AA488280	RC_AA403116 EC_H06397	RC W59961 s	RC_AA487449	RC_R43543	105614	RC_AA262179	RC_AA281451	RC_AA425691	KC_AA426376	KC_AA446000	774777 7047474	DC_A A 172056	C01169	AA075599	BC AA026356	X17059	1118321	9922X	RC AA464853	RC_AA227900	AA422025 s	RC AA346495	9755111	1174389	171090	PC H25577	S74445	DC AA419200	AA193297

4	3.9	3.8	3.7	3.6	3.6	3.5	3.5	3.4	3.4	3.3	3.3	3.3	3.2	5.2	3.1	1.5	5.1	n c	6.2	6.5	6.7	2.9	2.9	2.9	2.9	2.9	2.9 6.0	0.7	7.8 2.8	0.0	7.0 0.0	0.7	%. % o	8.7 0.0	2.7	2.7	2.7
Cytochrome B561	Homo sapiens mRNA for RB18A protein	ESTs Weakly similar to D9481.16 gene product	Human 60-kdal ribonucleoprotein (Ro) mRNA complete	ESTs	ESTs	NUCLEAR PORE GLYCOPROTEIN P62	ESTs	EST - L47276	Phosphoribosyl pyrophosphate amidotransferase	ESTs	ESTs	Homo sapiens DNA from chromosome 19p13.2 cosmids	Peroxisomal membrane protein 3 (35kD Zellweger	ESIS	Cyclin D1 (PRAD1 parathyroid adenomatosis 1)	SICE CONTRACTOR OF THE PROPERTY OF THE PROPERT	Homo sapiens ribonuclease P protein subunit p20 (KFF20)	ESIS	Homo sapiens M962 protein spliced isoform 2 mKNA	Thymidylate synthase	ESIS	ESTS	ESTs Weakly similar to coded for by C. elegans cDNA	ESTs Weakly similar to W02D9.2 [C.elegans]	DNA-BINDING PROTEIN NEFA PRECURSOR	H.sapiens mRNA for ras-related GTP-binding protein	ESIS	Homo sapiens forkhead protein (FKHKLI) mKNA	H.sapiens mRNA for M-phase phosphoprotein mpp3	ESTs Weakly similar to !!!! ALU SUBFAMILY J	ESI	FK506-binding protein 4 (39KU)	ESTs Weakly similar to elastin like protein	ESTS	Human lipid-activated protein kinase PRK2 mRNA	Human mRNA for platelet activating factor	ESTs
1179463	R44709	4 4 4 5 9 9 6 0	M25077	A 521240	N62873	A A 034365	791079	1.47276	A A 442070	A A 116036	R24237	AD000092	M86852	AA232939	X59798	W44928	AA401687	AA423827	N32333	D00596	AA399164	AA164293	AA203523	AA195936	AA485214	R50840	R97040	N21159	AA292765	AA480103	AA412497	H24460	R39234	R79617	U33052	D63391	W69160
1120462	077403 DV D44700	DC A 450060	N25077	0,000 M	KC_AA321240	KC_N03625	KC_AA034303	K/016/ 1.47276	07777	DC_AA116036	RC R24237 f	AD000092 cds	M86852	RC AA232939	X59798	RC W44928	RC AA401687	RC_AA423827	RC N32333	_D00596	RC AA399164	RC_AA164293	A A 203523	RC AA195936	RC_AA485214	RC R50840	RC_R97040	RC_N21159	RC AA292765	RC_AA480103	RC_AA412497	RC H24460 s	RC_R39234_r	RC R79617	1133052	1063391	RC_W69160

ESTs ESTs ESTs ESTs ESTs ESTs Aldehyde dehydrogenase 8 SRY (sex determining region Y)-box 4 Human mRNA for kinesin-related protein partial cds Androgen receptor (dihydrotestosterone receptor testicular ESTs Highly similar to SIGNAL RECOGNITION Human BRCA1-associated RING domain protein ESTs Highly similar to FK506-BINDING PROTEIN Homo sapiens mRNA for p115 complete cds ESTs Highly similar to GAG POLYPROTEIN [Avian Human mRNA for EBI1-ligand chemokine complete cds ESTs Weakly similar to Y53C12A.3 [C.elegans] ESTs Moderately similar to ALR [H.sapiens] Human mRNA for KIAA0310 gene complete cds MYB PROTO-ONCOGENE PROTEIN ESTs Weakly similar to PROBABLE UBIQUITIN Human IEF SSP 9502 mRNA complete cds KERATIN TYPE II CYTOSKELETAL 6D ESTs Weakly similar to F08G12.1 [C.elegans] Homo sapiens breast cancer putative transcription factor ESTs Moderately similar to !!!! ALU SUBFAMILY SQ SPLICING FACTOR U2AF 35 KD SUBUNIT H.sapiens mRNA for spermine synthase Human mRNA for transcriptional activator hSNF2b Human tetracycline transporter-like protein mRNA M23263 U37519 U76638 AA599219 W20391 W28362 4A43460 R81830 N49284 AA148516 X70683 AA609869 AA487202 AA476582 W80467 W80763 AA464013 M96982 Z49099 N92948 L00205 AA069285 N33063 AA250737 U77180 AA028028 AA485223 F02651 AA045481 AA598648 **AA130349** AB002308 AA449458 AA599674 AA447617 RC_AA609869 RC_W20391_s RC_AA028028 RC_AA485223 RC_N49284_s RC_AA069285 RC_AA599219 M96982 RC_N92948_s RC_AA148516 X70683 RC AA487202 RC_AA476582 RC_W80467 U37519 AA443460 U76638 RC_W80763 AA464013 RC Z40332 R81830 RC_N33063 RC_AA250737 W28362 RC_F02863 RC_AA130349 AB002308 RC_AA449458 RC_AA599674 RC F02651 RC_AA045481 Z49099 RC_AA598648 RC_AA447617 L00205

1.7	1.7	1.7	1.6	1.6	1.6	1.5	1.5	1.4	1.4	1.4	1 4		t: •	- -	1.4	1.3	1.3	1.3	13	7 7		7	7:1
ESTs	H.sanjens mRNA for 55.11 binding protein	ESTS Highly similar to C10 [H.sapiens]	Human mRNA for ceramide glucosyltransferase complete	EST. Westly similar to CLEAVAGE STIMULATION	ES 18 Wearth statutum to Caracia ESTs	ESTs	EST. Weakly similar to similar to mouse MMR1	H saniens HUNKI mRNA	ESTS	recentor associated	Human tumor necrosis ractor type a receptor agence	STONE SHOUL	SICO	ESIS	ESTS	ESTs Highly similar to Polio virus receptor protein	DI-N-ACETYLCHITOBIASE PRECURSOR	Homo saniens clone 23592 mRNA sequence	The state of the s	Homo sapiens agnii piecuisoi imata paitim cus	E013	Integral transmembrane protein i	ESIS
0037304 4	AA036366	1010101	AA26/071	040000	AA430437	AA026418	AA293508	AA219151	AA083918	AA150542	UI2595	AA262943	T16226	AA057193	AA171939	A A 101601	79759M	1012CM	AA+0/+72	AA156670	AA040696	L38961	U51698
	RC_AA056588	1518/U	RC_AA28/091	D50840	RC_AA456437	RC_AA026418	RC_AA293568	RC_AA279757	RC_AA085918	RC_AA156542	U12595	RC AA262943	RC T16226	DC 4 4057193	DC_A A 171939	DC_AA101601	100101AA JA	101C6TAT	RC_AA48/492	AA156670 s	$RC AA0406\overline{9}6$	L38961	1151698

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ratio tumor	vs breast	135.3	107.3	81.8	75.3	72.9	61.5	57.7	53.9	53.2	25	40.0	7.0.4	2.00		20.00	07	2, 20, 21, 22, 22, 23, 24, 24, 24, 24, 24, 24, 24, 24, 24, 24	23.0	6.62	0.22	20.3	1.7.	ָבַ עַ	- t	ק	2 5	<u> </u>	<u>.</u>		,	- 1	- 1	- (U
	Gene Name	FSTS	EST - RC AA453638	EST	EST - RC_AA461510	Collagen Type XI alpha 1	ESTs	EST - RC_N27351	H sapiens mRNA for Sm protein F	Human focal adhesion kinase (FAK) mRNA complete cds	ESTs Weakly similar to zinc-finger protein Zn72D [D.melanogaster]	ESTs	ESTs	ESTs	EST	EST	ESTs	ESTs	Inhibin beta A (activin A activin AB alpha polypeptide)	EST	ESTs	ESTs	ESTs	ESTs Highly similar to thyroid disease hypothetical autoantigen [H.saplens]	ESTs	Inhibin beta A (activin A activin AB alpha polypeptide)	ESTs	ESTs Weakly similar to B0334.4 [C.elegans]	Human fibroblast activation protein mRNA complete cds	S100 calcium-binding protein A7 (psoriasin 1)	H.sapiens Humig mRNA	Human melanoma antigen p15 mRNA complete cds	Homo sapiens mRNA for osteoblast specific factor 2 (OSF-20s)	Interferon (gamma)-induced cell line protein 10 from	Receptor protein-tyrosine kinase EDDR1	ESTs Highly similar to putative hydrophobic domain in amino acid positions 373	Human germline oligomeric matrix protein (COMP) mRNA complete cds
	Document	T70056	A A 4 5 3 5 3 8	AA461322	AA461510	R67275	AA453518	N27351	AA486737	AA453479	AA285050	AA291468	240805	AA169440	AA453641	AA609955	AA283905	AA412090	N27159	R65763	R97063	AA463189	AA505133	H83527	AA196768	X57579	AA191404	AA262969	AA436611	M86757	X72755	U19796	D13666	X02530	U48705	AA346385	L32137
	CI > 30	DC T70066	AC_173330	RC AA461322	RC AA461510	RC R67275 s	RC_AA453518	RC N27351	BC AA486737	RC AA453479 s	RC_AA285050_s	RC_AA291468	RC_Z40805	RC_AA169440	RC_AA453641	RC_AA609955	RC_AA283905	RC_AA412090	RC_N27159_s	RC_R65763	RC_R97063	RC_AA463189	RC_AA505133	H83527_s	RC_AA196768_s	X57579	RC_AA191404	RC_AA262969_f	RC AA436611_s	M86757	X72755	U19796	D13666	X02530	U48705 rna1	RC AA346385	L32137

211	YES	YES	YES	YES	YES	YES	YES	YES) 	ר ב ר ב	у П М	2 L2	S II X	2
4.4	4.2	3.2	2.9	2.9	2.8	2.7	2.6	3.6		7 7	- -	o u		7.1
,	ES1S	remain Similar to COATOMER GAMMA SUBUNIT ISaccharomyces cere	ESTS rightly similar to coded for by C. elegans cDNA vk10c10.3 [C.elegans]	ES IS Vyčasky Silliliai to cooca to 3) of coosa to 2)	n.saplens more of sacratica (control of sacratica (control of sacratica of sacratica (control of sacratica of	Trybus-billioning protein 4 (2000)	Edits Weakly similar to roof this of the constant	ESTs Weakly similar to semaphorin C (M. musculus)	ESTs	ESTs	Glucose-6-phosphate dehydrogenase	ESTs Weakly similar to monocytic leukaemia zinc finger protein [H.Sapiens]	ESTs	Human mRNA for Tob complete cds
	AA422025	H25577	T15916	AA203523	R50840	H24460	AA125969	AA293300	AA447617	R50333	X55448	AA443962	AA293568	D20342
	AA422025_s	RC_H25577	RC_T15916	AA203523	RC_R50840	RC_H24460_s	RC_AA125969	RC_AA293300_s	RC_AA447617	RC R50333 I	X55448 cds1	RC AA443962	RC_AA293568	RC_D20342_

Affymetrix ID	Accession	Gene Name	ratio tumor vs breast	ORF structural info
RC_T79956	T79956	ESTs	135 3	?
RC_AA453640	AA453640	ESTs	121.4	other
RC_AA453638	AA453638	EST - RC_AA453638	107 3	3
RC_AA461322	AA461322	EST	81.8	?
RC_AA461510	AA461510	EST - RC_AA461510	75.3	other
RC_R67275_s	R67275	Collagen type XI alpha 1	72.9	other
RC_AA453518	AA453518	ESTs	61 5	other
RC_N27351	N27351	EST - RC_N27351	57.1	?
RC_AA486737	AA486737	H.sapiens mRNA for Sm protein F	53 9	TM
RC_AA453479_s	AA453479	Human focal adhesion kinase (FAK) mRNA complete cds	53.2	other
RC_AA285050_s	AA285050 AA291468	ESTs Weakly similar to zinc-finger protein Zn72D [D.melanogaster]	52	other
RC_AA291468 RC_Z40805	Z40805	ESTs ESTs	46.8	TM
RC_AA169440	AA169440	ESTs	45.7	other
D90041_s	D90041	ARYLAMINE N-ACETYLTRANSFERASE MONOMORPHIC	38.9 33.6	other ?
RC_AA621202	AA621202	ESTs	33.5	r other
RC_AA232294	AA232294	EST - RC_AA232294	32.6	other
RC_R86839	R86839	EST - RC_R86839	32.4	other
\$70585_ma1	S70585	GLYCOPROTEIN HORMONES ALPHA CHAIN PRECURSOR	31.3	ت مرابط ت
RC_AA453641	AA453641	EST	31.1	ss.
RC_AA609955	AA609955	EST	30.6	TM
RC_AA283905	AA283905	ESTs	28.3	?
RC_AA211831	AA211831	EST - RC_AA211831	28.1	TM
RC_AA412090	AA412090	ESTS	28	other
RC_AA421289	AA421289	ESTs Weakly similar to ZINC FINGER PROTEIN MFG1 [Mus musculus]	25.5	other
RC_N27159_s	N27159	Inhibin beta A (activin A activin AB alpha polypeptide)	25.5	other
RC_T16687	T16687	ESTs	25 1	other
RC_R65763	R65763	EST	23.9	?
RC_AA487987	AA487987	EST	23 8	ТМ
RC_H99309	H99309	Human TFIID subunits TAF20 and TAF15 mRNA complete cds	23 6	other
RC_R97063	R97063	ESTs	22 8	other
RC_AA232940	AA232940	EST - RC_AA232940	21.7	other
RC_AA463189	AA463189	ESTs	20 9	TM
RC_AA421171	AA421171	ESTs	195	other
RC_AA251875_f	AA251875	ESTs Moderately similar to POL POLYPROTEIN [Feline endogenous virus ece1]	19 4	other
RC_AA054228	AA054228	ESTs	17.7	other
RC_D51215_f	D51215	EST - RC_D51215_f .	17 4	other
RC_AA505133	AA505133	ESTs	17 1	other
RC_AA621462	AA621462	CARCINOEMBRYONIC ANTIGEN PRECURSOR	17 1	other
RC_AA232508	AA232508	ESTs	17	other
RC_AA024659_f	AA024659	H sapiens mRNA for hHKb1 protein	16 9	other
RC_AA211158	AA211158	EST - RC_AA211158	16 8	other
RC_AA488191	AA488191	ESTs	168	other
RC_AA290674_s	AA290674	Human 4E-binding protein 1 mRNA complete cds	16 3	other
RC_AA481883	AA481883	ESTs	16 2	TM
RC_AA196721	AA196721	EST - RC_AA196721	16 1	other
RC_AA196768_s	AA196768	ESTs	16 1	other
H83527_s	H83527	ESTs Highly similar to thyroid disease hypothetical autoantigen [H sapiens]	16 1	other
RC_D51172	D51172	ESTs	15 9	other
RC_T25875	T25875	Homo sapiens clone 23967 unknown mRNA partial cds	15.8	other
X57579 J05068	X57579 J05068	Inhibin beta A (activin A activin AB alpha polypeptide)	15.8	7
RC_AA487233	AA487233	TRANSCOBALAMIN I PRECURSOR	15.7	SS.
RC_AA479969		ESTs Moderately similar to !!!! ALU SUBFAMILY SP WARNING ENTRY !!!! [H.sapiens]	156	TM
RC_AA191404	AA479969 AA191404	ESTs ESTs	15.4	other
RC_AA262969_f	AA262969	ESTs Weakly similar to B0334 4 [C.elegans]	152	other
RC_AA436611_s	AA436611	Human fibroblast activation protein mRNA complete cds	14 9	
RC_AA250843_s	AA250843	Interferon regulatory factor 5	147	SS.
RC_AA461297	AA461297	ESTs	14 6 14 6	
RC_R51309	R51309	ESTs	146	other other
RC_AA280679	AA280679	ESTs	14.4	omer 7
RC_AA412029	AA412029	ESTs	14.4	other
RC_AA430032	AA430032	ESTs Moderately similar to PTTG gene product [R norvegicus]	14 4	7

FIGURE 7
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M81057	M81057	Carboxypeptidase B1 (tissue)	14 4	SS.
RC_R07976	R07976	ESTs Highly similar to HYPOTHETICAL 21.5 KD PROTEIN CO8B11 9 IN	14 3	7
_		CHROMOSOME II (Caenorhabditis elegans)		
U75285_ma1	U75285	Human effector cell protease receptor-1 (EPR-1) gene partial cds	143	?
RC_R46627	R46627	ESTs Charmonica A (coverburgid population 4)	14.2	other
RC_AA461559	AA461559	Chromogranin A (parathyroid secretory protein 1)	14.1	?
AA092129_f	AA092129	ESTs Moderately similar to 25E8.I [D.melanogaster]	14	other
RC_AA436893	AA436893	ESTs Weakly similar to TH1 protein [D melanogaster]	14	?
RC_AA465345	AA465345	ESTs	13 9	other
RC_AA486538	AA486538	ESTs Weakly similar to hypothetical protein 1 [H.sapiens]	13.9	other
M23263	M23263	Androgen receptor (dihydrotestosterone receptor testicular feminization spinal and bulbar muscular atrophy Kennedy disease)	13.9	TM
RC_D20379	D20379	ESTs	13 7	other
RC_AA076138	AA076138	Homo sapiens histone macroH2A1.2 mRNA complete cds	13.5	other
RC_W60486	W60486	ESTs Moderately similar to T11G6.8 [C.elegans]	13.5	other
RC_AA032243	AA032243	EST - RC_AA032243	13 4	other
RC_AA045074_s	AA045074	ESTs Weakly similar to 52-kD SS-A/Ro autoantigen [H sapiens]	13.4	other
RC_F01444_f	F01444	Homo sapiens KIAA0440 mRNA partial cds	13 4	other
AA401334	AA401334	ESTs	13.3	other
RC_D60354_s	D60354	Human mRNA for KIAA0007 gene partial cds	13.3	other
RC_T78922_s	178922	Homo sapiens mRNA for SCGF-beta complete cds	13.3	other
RC_AA406635	AA406635	ESTs	13.1	other
RC_AA412065	AA412065	EST - RC_AA412065	13.1	other
RC_AA431350	AA431350	ESTs Moderately similar to !!!! ALU SUBFAMILY SC WARNING ENTRY !!!! [H.sapiens]	13.1	other
RC_AA431738	AA431738	EST	13.1	?
RC_R61740_f	R61740	Homo sapiens regulator of G-protein signalling 12 (RGS12) mRNA complete cds	13	other
RC_R54950	R54950	ESTs	128	other
RC_AA405488	AA405488	ESTs	12.7	TM
RC_AA418749	AA418749	EST	12 7	other
AA037285	AA037285	Homo sapiens mRNA for A+U-rich element RNA binding factor complete cds	12 5	other
RC_AA233796	AA233796	ESTs	12 5	other
RC_AA219305	AA219305	EST	124	2
RC_AA252245	AA252245	ESTs	12 4	TM
RC_AA041276	AA041276	ESTs Weakly similar to "" ALU SUBFAMILY SX WARNING ENTRY " [H sapiens]	12 3	7
RC_AA463874	AA463B74	Homo sapiens ES/130 mRNA complete cds	123	TM
RC_AA099404_s	AA099404	ESTs	12 2	other
RC_AA443985	AA443985	ESTs	12 2	2
RC_AA461528	AA461528	ESTs	122	TM
RC_AA214305	AA214305	ESTs	12.1	other
AA220223	AA220223	Fibroblast growth factor receptor 2 (bacteria-expressed kinase keratinocyte growth factor receptor craniofacial dysostosis 1 Crouzon syndrome Pfeiffer syndrome Jackson-Weiss	12.1	SS,TM
RC_AA478571	AA478571	syndrome)	12 1	The
U31875	U31875	Glutamine-fructose-6-phosphate transaminase	12 1	TM
RC_AA253217	AA253217	Human Hep27 protein mRNA complete cds ESTs	11 8	TM
RC_AA470074	AA470074	ESTs	115	other
RC_AA236010	AA236010	ESTs	11.4	other
RC_AA430002	AA430002	ESTs	11.4	other other
D82307	D82307	ESTs Weakly similar to TH1 protein [D melanogaster]	11.4	other
J03589	J03589	UBIQUITIN-LIKE PROTEIN GDX	11 4	2
RC_AA179298 RC_R22952_s	AA179298	Homo sapiens chromosome 9 P1 clone 11659	11 3	other
	R22952 W56363	ESTs	11 3	7
RC_W56363		ESTs Weakly similar to extracellular protein [H sapiens]	11 3	TM
RC_AA449232	AA449232	EST	11.2	?
RC_AA444054	AA444054	ESTs Weakly similar to transmembrane protein [H sapiens]	11.1	?
RC_AA281733	AA281733	ESTs	11	other
RC_AA452601	AA452601	EST	11	7
RC_AA035630	AA035630	Homo sapiens U4/U6 small nuclear ribonucleoprotein hPrp4 mRNA complete cds	109	other
RC_AA235117	AA235117	ESTs Weakly similar to espin (R norvegicus)	109	other
RC_AA279418	AA279418	ESTs	109	TM
RC_AA432069	AA432069	ESTs	108	?
RC_AA453630	AA453630	EST	10.7	?
RC_W44657	W44657	EST	10 7	7
RC_AA405098	AA405098	ESTs Weakly similar to MOESIN/EZRIN/RADIXIN HOMOLOG [D melanogaster]	106	other
RC_R40431	R40431	ESTs	106	other
RC_AA411425	AA411425	ESTs	10 5	other

FIGURE 7 (cont.)
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RC_AA423956	AA423956	ESTs	10.5	Othor
RC_AA253170	AA253170	EST	10.5	other ?
RC_AA459347	AA459347	ESTs	10.4	other
RC_AA599259_s	AA599259	Human splicesomal protein (SAP 61) mRNA complete cds	10.4	other
X62078	X62078	GANGLIOSIDE GM2 ACTIVATOR PRECURSOR	10.4	SS,
RC_AA251430	AA251430	ESTs Highly similar to RAS-RELATED PROTEIN RAB-10 [Canis familiaris]	103	other
RC_AA470156	AA470156	ESTs Weakly similar to dynein 74K chain cytosolic [R norvegicus]	103	SS,
RC_T64933_f	T64933	ESTs	103	other
RC_AA280609	AA280609	ESTs Weakly similar to K02B2.3 gene product [C.elegans]	10.2	other
RC_AA281290	AA281290	ESTs Highly similar to ZINC FINGER PROTEIN 85 [Homo sapiens]	10.2	?
RC_AA449832	AA449832	ESTs	10.1	other
RC_AA427898	AA427898	ESTs Weakly similar to trabecular meshwork inducible glucocorticoid response protein	10	other
RC_AA609867	AA609867	[H.sapiens] ESTs Weakly similar to No definition line found [C.elegans]	10	other
RC_AA465158	AA465158	EST	9.9	?
RC_R49198_i	R49198	H.sapiens DAP-3 mRNA	9.9	?
RC_AA112396	AA112396	ESTS	9.8	other
RC_AA207015	AA207015	ESTs	9.8	other
RC_AA228030	AA228030	ESTs	9.7	TM
RC_AA447982	AA447982	ESTs Weakly similar to R01H10 6 [C.elegans]	9.7	other
M34338	M34338	Spermidine synthase	9.7	other
RC_R06986_f	R06986	ESTs	9.7	?
Z14982_ma1	Z14982	PROTEASOME COMPONENT C13 PRECURSOR	9.7	?
RC_AA176247	AA176247	EST	9.6	other
RC_T97341	T97341	EST - RC_T97341	9.6	?
W26392	W26392	ESTs Highly similar to OVOSTATIN PRECURSOR [Gallus gallus]	9.6	other
RC_AA143190_s	AA143190	ESTs Highly similar to HYPOTHETICAL 23.1 KD PROTEIN IN SHP1-SEC17	95	TM
RC_AA452578	AA452578	INTERGENIC REGION [Saccharomyces cerevisiae] ESTs	9.5	other
RC_AA258057	AA258057	ESTs	9.4	other
RC_AA282914	AA282914	ESTs	9.4	other
RC_AA461476	AA461476	ESTs Highly similar to PUTATIVE ATP-DEPENDENT RNA HELICASE C31A2 07C	9.4	other
RC_W87751	W87751	[Schizosaccharomyces pombe] ESTs	9.4	
RC_W92713	W92713	ESTs	9.4	other other
RC_AA262111	AA262111	ESTs	93	other
RC_AA490929	AA490929	EST	93	?
RC_N21678	N21678	EŞTS	93	?
RC_N70690	N70690	ESTs	93	other
RC_N80716	N80716	ESTs	93	other
RC_AA007344	AA007344	ESTs	92	other
D14657	D14657	Human mRNA for KIAA0101 gene complete cds	92	other
RC_W73140	W73140	ESTs Highly similar to TRYPSINOGEN ANIONIC PRECURSOR [Canis familiaris]	92	other
RC_AA243020	AA243020	H.sapiens mRNA for disintegrin-metalloprotease (partial)	91	SS,TM
RC_AA431478	AA431478	ESTs	9 1	other
RC_AA447666_s	AA447666	Human CENP-F kinetochore protein mRNA complete cds	9.1	other
RC_T16308_f	T16308	ESTs	91	other
RC_R38919_i	R38919	EST	9	other
RC_R60223_s	R60223	ESTs	9	other
RC_R70379_s	R70379	Human germline IgD chain gene C-region C-delta-1 domain	9	2
HG2981-HT3127	HT3127	EST - HG2981-HT3127	9	?
M86757	M86757	S100 calcium-binding protein A7 (psoriasin 1)	89	SS,TM
RC_AA347209_s	AA347209	Human mRNA for KIAA0324 gene partial cds	88	other
RC_AA485041	AA485041	ESTs	8.8	other
X72755	X72755	H sapiens Humig mRNA	8.8	TM
RC_AA443342_s	AA443342	ESTs	8.7	other
RC_AA481281	AA481281	ESTs	8.7	other
RC_T96361_s	T96361	MULTIFUNCTIONAL AMINOACYL-TRNA SYNTHETASE	87	other
RC_AA608723	AA608723	ESTs	86	other
RC_H18027_s M86752	H18027 M86752	Homo sapiens clone 23785 mRNA sequence TRANSCORMATION SENSITIVE PROTEIN IEE SSR 2524	8.6	SS,
RC_AA113011_s	AA113011	TRANSFORMATION-SENSITIVE PROTEIN IEF SSP 3521 Human mRNA for KIAA0314 gene partial cds	86 86	other
RC_AA457018	AA457018	ESTs	8 5 8 5	other SS,
RC_H96237_s	H96237	Collagen type XI alpha 1	85	აა, other
RC_AA024835	AA024835	Homo sapiens Shab-related delayed-rectifier K+ channel alpha subunit (KCNS3) mRNA	84	TM
_	-	complete cds		1,41
		FIGURE 5 ()		

FIGURE 7 (cont.)
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	RC_N99976	N99976	ESTs	8 4	other
	\$85655	S85655	Prohibitin	8.4	other
	RC_T65004	T65004	EST - RC_T65004	8 4	other
	RC_AA489510_s	'AA489510	Homo sapiens clone 23716 mRNA sequence	83	7
	RC_H72948_s	H72948	ESTs Highly similar to BONE/CARTILAGE PROTEOGLYCAN I [Bos taurus]	8.3	SS,
	RC_N93197	N93197	ESTs	8.3	other
	RC_Z39971_s	Z39971	ESTs	83	other
	RC_AA236037	AA236037	ESTs Highly similar to HYPOTHETICAL 37.8 KD PROTEIN 80285 4 IN CHROMOSOME III [Caenorhabddis elegans]	8.2	7
	J05070	J05070	Matrix metalloproteinase 2 (gelatinase A collagenase type (V)	8.2	TM
	RC_AA419225	AA419225	Human mariner-like element-containing mRNA clone pcHMT1	8.1	other
	RC_D60302	D60302	ESTs	8.1	other
	RC_H98621_s	H98621	Homo sapiens mRNA for KIAA0617 protein complete cds	8.1	?
	RC_R40177	R40177	ESTs	8.1	other
	RC_AA233545 RC_AA436370	AA233545 AA436370	ESTS Weakly similar to HYPOTHETICAL 26.1 KD PROTEIN IN RIB5-SHM1 INTERGENIC REGION [Saccharomyces cerevisiae] ESTS Highly similar to ADP-RIBOSYLATION FACTOR-LIKE PROTEIN 4 [Rattus	8	TM other
	_		norvegicus)		
	RC_F01538_s	F01538	RAP1 GTPase activating protein 1	8	other
٠	RC_N39415	N39415	ESTs Highly similar to OSTEOINDUCTIVE FACTOR PRECURSOR [Bos taurus]	8	SS,
	RC_W93659	W93659	ESTs EGT-	8	other
	RC_AA053319	AA053319	ESTs ESTs	7.9	TM
	RC_AA235009	AA235009	EST	7.9	?
	RC_AA421750 RC_AA447574	AA421750 AA447574	EST's	7.9	TM
		AA458882	ESTs Weakly similar to LINE-1 REVERSE TRANSCRIPTASE HOMOLOG [Homo	7.9 7.9	other
	RC_AA458882 RC_N33011_s	N33011	Replication protein A (E coli RecA homolog RAD51 homolog)	7.9	other
	RC_N53950	N53950	EST - RC_N53950	7.9	other
	RC_N69464	N69464	ESTs	7.9	other
	RC_R01634	R01634	ESTs	7.9	other
	RC_R47948_i	R47948	ESTs	79	other
	RC_AA150182	AA150182	ESTs Weakly similar to HYPOTHETICAL 88 1 KD PROTEIN K02D10.1 IN	7.8	other
	RC_AA342084	AA342084	CHROMOSOME III [C.elegans] EST - RC_AA342084	7.8	other
	RC_AA417213	AA417213	ESTs	7.8	other
	RC_AA446486	AA446486	Homo sapiens Ran binding protein 2 (RanBP2alpha) mRNA partial cds	78	٤
	RC_AA609170	AA609170	EST	78	?
	RC_N46435	N46435	EST - RC_N46435	7.8	other
	RC_N54916	N54916	Human mRNA for KiAA0136 gene partial cds	78	other
	RC_AA459389	AA459389	Homo sapiens mRNA for tyrosyl sulfotransferase-2	77	TM
	RC_AA463693	AA463693	ESTs .	7.7	other
	RC_T88814	T88814	ESTs	77	TM
	RC_AA446008	AA44600B	EST	76	7
	RC_H99879	H99879	ESTs Highly similar to EPIDERMAL GROWTH FACTOR PRECURSOR [Mus musculus]	76	other
	RC_T03306	T03306	Homo sapiens clone 24703 beta-tubulin mRNA complete cds	76	3
	RC_T68871	T68871 TIGR - HG2981-	ESTs	76	other
	HG2981-HT3938	HT3938	EST - HG2981-HT3938	76	7
	U19796	U19796	Human melanoma antigen p15 mRNA complete cds	76	other
	AA094752	AA094752	Calcineurin B	75	other
	RC_AA282074	AA282074	ESTs	75	other
	RC_AA442767 RC_AA454566	AA442767 AA454566	Tyrosine 3-monooxygenase/tryptophan 5-monooxygenase activation protein beta polypeptide Human mRNA for KIAA0170 gene complete cds	75 75	other ?
	RC_AA476937_s	AA476937	ESTs	75 75	
	D13666	D13666	Homo sapiens mRNA for osteoblast specific factor 2 (OSF-2os)	75 75	other SS.
	RC_N67119	N67119	ESTs	75	33. ?
	W01296	W01296	EST - W01296	7.5	TM
	RC_AA069476_s	AA069476	H sapiens mRNA for surface glycoprotein	7.4	other
	RC_AA287061	AA287061	ESTs	7.4 7.4	other
	RC_AA410190	AA410190	ESTs	7.4	other
	RC_AA411952	AA411952	Homo sapiens mRNA for GalT4 protein	7.4	7
	RC_AA412477	AA412477	EST	7.4	2
	RC_AA459392	AA459392	ESTs	7.4	ather
	RC_AA486256	AA486256	ESTs Moderately similar to breast cancer suppressor element Ishmael Upper RP2 [H.sapiens]	7 4	7
	RC_AA599042	AA599042	EST	7 4	7
	RC_AA609309	AA609309	ESTs	74	other

FIGURE 7 (cont.)
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RC_D59489	D59489	ESTs	7.4	SS,TM
RC_N67889	N67889	ESTs	7.4	other
RC_W73520	W73520	ESTs Highly similar to HYPOTHETICAL 28.5 KD PROTEIN ZK1236.7 IN CHROMOSOME III [Caenorhabditis elegans]	7.4	other
X02530	X02530	Interferon (gamma)-induced cell line protein 10 from	7.4	SS,
RC_AA283006	AA283006	ESTs Highly similar to CHROMOSOME CONDENSATION PROTEIN DPY-27 [Caenorhabditis elegans]	7.3	other
RC_AA426372_s	AA426372	Human mRNA for histone H1x complete cds	7.3	other
RC_AA443794	AA443794	ESTs	73	other
RC_AA446869	AA446869	ESTs	7.3	TM
RC_F13642	F13642	ESTs	7.3	other
RC_N21321_i	N21321	ESTs	7.3	?
RC_AA088458	AA088458	ESTs Weakly similar to !!!! ALU SUBFAMILY J WARNING ENTRY !!!! [H.sapiens]	7.2	other
RC_AA121315	AA121315	ESTs	7.2	other
RC_AA234921	AA234921	ESTs	7.2	other
RC_AA427950	AA427950	EST - RC_AA427950	72	?
RC_AA432130	AA432130	ESTs Moderately similar to !!!! ALU SUBFAMILY SX WARNING ENTRY !!!! [H.sapiens]	7.2	?
RC_R65593_s	R65593	Homo sapiens mRNA for kynurenine 3-monooxygenase	7.2	TM
RC_AA236177	AA236177	ESTs	7.1	7
RC_AA258482_s	AA258482	Human mRNA for zinc finger protein complete cds	7.1	SS.
RC_AA282143_s	AA282143	H.sapiens mRNA for melanoma growth regulatory protein MIA	7.1	SS,
RC_AA283003	AA283003	ESTs	7.1	other
RC_AA287870_s	AA287870	Lymphotoxin-beta	7.1	other
AA310967_s	AA310967	ESTs Weakly similar to T04A8.11 [C elegans]	7.1	other
RC_AA410373	AA410373	ESTs	7.1	າ
RC_AA421158	AA421158	ESTs	7.1	SS,
C00225_s	C00225	ESTs Highly similar to HYPOTHETICAL 52.8 KD PROTEIN T05E11 5 IN	7.1	TM
DO 110004 6	E42004	CHROMOSOME IV [Caenorhabditis elegans]		
RC_F13694_f	F13694	ESTs DA MONTO	71	7
RC_N29431	N29431	EST - RC_N29431	7.1	?
RC_N67239	N67239	ESTs	7.1	other
RC_T10082_f	T10082	ESTs	7.1	other
RC_AA310499	AA310499	ESTs	7	other
RC_AA449351	AA449351	ESTs Weakly similar to similar to deoxyribose-phosphate aldolase [C elegans]	7	other
RC_D57389_f	D57389	EST	7	other
RC_N34686	N34686	Homo sapiens clone 23915 mRNA sequence	7	7
RC_Z40345	Z40345	ESTs Weakly similar to T06D8 5 (C elegans)	7	TM
RC_AA410441	AA410441	ESTs	69	other
RC_AA505093	AA505093	ESTs	69	other
RC_N71704	N71704	ESTs	69	other
U48705_ma1	U48705	Receptor protein-tyrosine kinase EDDR1	. 69	2
RC_AA127818_i	AA127818	ESTs	68	7
RC_AA346385 RC_AA411204	AA346385 AA411204	ESTs Highly similar to putative hydrophobic domain in amino acid positions 373-390 [H.sapiens] ESTs	68	other
RC_AA416876			68	TM.
RC_AA419461	AA416876 AA419461	ESTs Weakly similar to TRANSFORMATION-SENSITIVE PROTEIN IEF SSP 3521 [H.sapiens]	68	other
RC_AA446966	AA446966	ESTS	68	other
RC_AA496569	AA496569	EST Highly significants MAN MI TOMA CONTRICTAGE (Company)	68	other
-	D51229	ESTs Highly similar to VALYL-TRNA SYNTHETASE [Fugu rubripes]	68	other
RC_D51229_f RC_F02254_s	F02254	Human clone 23589 mRNA sequence	6.8	TM
RC_H18428_s	H18428	H sapiens mRNA for FAST kinase ESTs Weakly similar to "!!! ALU SUBFAMILY J WARNING ENTRY !!" [H sapiens]	68	other
RC_AA034069	AA034069	ESTS	68	
RC_AA127058	AA127058	ESTs	67	other
RC_AA435849	AA435849		67	TM
RC_H99935_s	H99935	ESTs Moderately similar to unknown protein [H sapiens]	67	other
L27841	L27841	Interfeukin 6 signal transducer (gp130 oncostatin M receptor) Human autoantigen pericentriol material 1 (PCM-1) mRNA complete cds	67	TM
M11718	M11718	Collagen type V alpha	67	other
RC_N50550			67	
RC_T92935	N50550 T92935	Homo sapiens mRNA for Efs1 complete cds	67	other
U24169	U24169	ESTS	67	other
U30246	U30246	Human JTV-1 (JTV-1) mRNA complete cds	67	other
RC_AA098834_s	AA098834	Human burnetanide-sensitive Na-K-Cl cotransporter (NKCC1) mRNA complete cds	67	TM
RC_AA283198	AA283198	Nuclear factor of kappa light polypeptide gene enhancer in B-cells 1 (p105) ESTs	66	TM
RC_AA421782	AA421782	ESTs	66	other
RC_AA505141	AA505141	ESTs	66	other
			66	other

FIGURE 7 (cont.)
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DC D60341	D60341	ESTs	66	SS.
RC_D60341 RC_N26904	N26904	ESTS Highly similar to FK506-BINDING PROTEIN PRECURSOR [Mus musculus]	66	TM
RC_R40606	R40606	ESTs Highly similar to SKD3 [M musculus]	66	other
RC_R51988	R51988	ESTs	6.6	other
RC_T03790	T03790	ESTs	66	other
RC_W72455	W72455	ESTs	66	TM
RC_AA100364	AA100364	ESTs	6.5	other
AA236384	AA236384	ESTs Highly similar to COP1 REGULATORY PROTEIN (Arabidopsis thaliana)	6.5	other
RC_AA431085	AA431085	EST	6.5	?
RC_AA446591	AA446591	ESTs	65	other
RC_R06700	R06700	ESTs	6.5	other
W49521	W49521	Human prolyl 4-hydroxylase alpha (II) subunit mRNA complete cds	6.5	?
RC_AA195651	AA195651	EST	6.4	?
RC_AA430211	AA430211	ESTs	6.4	other
RC_T15991	T15991	ESTs	6.4	TM
RC_T17119	T17119	ESTs	6.4	TM
U89606	U89606	Human pyridoxal kinase mRNA complete cds	6.4	other
X54326	X54326	MULTIFUNCTIONAL AMINOACYL-TRNA SYNTHETASE	6.4	other
RC_AA157814	AA157814	ESTs	6.3	other
RC_AA443658	AA443658	Homo sapiens lamin 8 receptor homolog TM7SF2 (TM7SF2) mRNA complete cds	6.3	TM
RC_AA621169	AA621169	ESTs	6.3	other
RC_D20168	D20168	Human mRNA for KIAA0050 gene complete cds	6.3	other
D81608	D81608	H.sapiens mRNA for RNA polymerase II subunit	6.3	TM
RC_H57330	H57330	EST	63	2
L77701	L77701	Homo sapiens COX17 mRNA complete cds	63	other
RC_N48166	N48166	ESTs	63	other
RC_R65826	R65826	Homo sapiens mRNA for KIAA0549 protein partial cds	63	SS.
W19662	W19662	ESTs CONTRACTOR CONTRA	63	other
X70649	X70649	Homo sapiens DDX1 gene complete CDS	6.3 6.2	?
RC_AA024664_s	AA024664	Human NADH ubiquinone oxidoreductase subunit B13 (B13) mRNA complete cds	6.2	other
RC_AA098874	AA098874	ESTs	6.2	other
RC_AA152178	AA152178	ESTs ESTs	6.2	other other
RC_AA279943	AA279943 AA412106	ESTs	6.2	other
RC_AA412106 RC_AA621721	AA621721	ESTs	62	other
RC_N38959_f	N38959	Homo sapiens chaperonin containing t-complex polypeptide 1 beta subunit (Cctb) mRNA	62	other
		complete cds		
U40271	U40271	Protein-tyrosine kınase 7	62	TM ?
X52150_rna1_s	X52150	Arylsulfatase A	62	
X86018	X86018	H sapiens mRNA for MUF1 protein	62 61	other ?
RC_AA133199	AA133199	ESTs	61	other
RC_AA398740 RC_AA405505	AA398740 AA405505	ESTs Homo sapiens mRNA for putative RNA helicase 3' end	61	other
RC_AA416568	AA416568	ESTs	61	other
RC_AA448349	AA448349	ESTs	61	2
AA455331	AA455331	ESTs	61	TM
RC_AA479933_f	AA479933	ESTs	61	other
RC_AA521080	AA521080	ESTs	61	other
RC_AA600257_s	AA600257	ERGIC-53 PROTEIN PRECURSOR	6 1	TM
RC_D20280	D20280	ESTs	61	other
RC_H55748	H55748	ESTs	61	other
RC_H55915	H55915	ESTs Weakly similar to LINE-1 REVERSE TRANSCRIPTASE HOMOLOG [H sapiens]	61	?
L32137	L32137	Human germline oligomeric matrix protein (COMP) mRNA complete cds	61	other
L37747_s	L37747	LAMIN B1	61	7
RC_N92593	N92593	ESTs	61	SS,
RC_R99978	R99978	ESTs Weakly similar to line-1 protein ORF2 (H sapiens)	61	?
RC_T77733_s	T77733	Tubulin gamma polypeptide	6.1	?
U78525	U78525	Human eukaryotic translation initiation factor (eIF3) mRNA complete cds	6 1	other
RC_W69807	W69807	ESTs Highly similar to GOLIATH PROTEIN [Drosophila melanogaster]	61	other
RC_AA406137	AA406137	EST	6	?
RC_AA521103	AA521103	ESTs	6	other
RC_AA609277	AA609277	ESTs	6	other
J04177	J04177	Collagen type XI alpha 1	6	olher
RC_R33663_s	R33663	ESTs	6	7

FIGURE 7 (cont.)
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RC_T16660	T16660	ESTs	6	other
U73514	U73514	Homo sapiens short chain L-3-hydroxyacyl-CoA dehydrogenase (SCHAD) mRNA complete cds	6	other
RC_AA223730	AA223730	ESTs	5.9	other
RC_AA292655	AA292655	ESTs	59	other
RC_N26391	N26391	ESTs	59	other
RC_N50744	N50744	ESTs	59	other
RC_T88953	T88953	ESTs	59	TM
HG3748-HT4018		EST - HG3748-HT4018	5.9	?
RC_W38407	HT4018 W38407	ESTs	5.9	other
RC_W63563_s	W63563	Homo sapiens scaffold attachment factor B (SAF-B) mRNA partial cds	5.9	7
RC_Z41619_\$	Z41619	ESTs Weakly similar to keratin 8 type II cytoskeletal embryonic [M.musculus]	5.9	other
RC_AA279292	AA279292	ESTs	5.8	other
RC_AA287665	AA287665	ESTs	58	other
RC_AA422007	AA422007	ESTs	58	TM
RC_AA425379	AA425379	ESTs	58	TM
RC_AA427925_s	AA427925	ESTs Weakly similar to PROCOLLAGEN ALPHA 1(II) CHAIN PRECURSOR [Homo	58	?
RC_AA430673	AA430673	sapiens) ESTs	58	other
RC_AA441801	AA441801	ESTs	5.8	other
RC_AA463740_s	AA463740	ESTs	5.8	other
RC_H89987_s	H89987	Human multidrug resistance-associated protein homolog (MRP5) mRNA partial cds	58	TM
RC_H94843	H94843	ESTs	58	other
M25753	M25753	Cyclin B1	58	other
RC_N80183	N80183	ESTs	58	other
RC_T67463_s	T67463	CATHEPSIN K PRECURSOR	58	other
U65932	U65932	Human extracellular matrix protein 1 (ECM1) mRNA complete cds	58	?
RC_AA149624	AA149624	Homo sapiens mRNA for follistain-related protein (FRP) complete cds	57	TM
RC_AA192334	AA192334	ESTs	57	other
RC_AA207105	AA207105	EST	57	?
RC_AA442763	AA442763			
-		ESTs Highly similar to G2/MITOTIC-SPECIFIC CYCLIN B2 [Mesocricetus auratus] ESTs	57	other
AA443251	AA443251	ESTs .	57	other
RC_AA454562	AA454562		57	other
RC_AA459945	AA459945	Homo sapiens mRNA for KIAA0585 protein partial cds	57	other
RC_AA478794	AA478794	ESTs	57	other
RC_AA609473	AA609473	ESTs	5.7	other
RC_F09058	F09058	ESTs	57	other
RC_H54430	H54430	ESTs	57	other
M34677	M34677	FACTOR VIII INTRON 22 PROTEIN	5 7	?
RC_N27563	N27563	ESTs Standard	57	other
RC_R02572	R02572	Fibronedin 1	57	other
RC_R09166	R09166	ESTs	57	2
RC_R85829	R85829	EST	5 7	other
HG2981-HT3125	HT3125	EST - HG2981-HT3125	5 7	7
U56402	U56402	Homo sapiens clone 24522 mRNA sequence	57	other
W30943	W30943	ESTs	57	other
RC_AA040154	AA040154	ESTs	56	other
AA116095	AA116095	ESTs Weakly similar to T12D8 / [C elegans]	56	other
RC_AA147884	AA147884	ESTs	56	TM
RC_AA149754_i	AA149754	EST	56	7
RC_AA232956	AA232956	E\$Ts	56	other
RC_AA397919	AA397919	ESTs	56	other
RC_AA398212	AA398212	ESTs	56	other
RC_AA398264	AA398264	Homo sapiens clone 23736 mRNA sequence	56	other
RC_AA406169	AA406169	Homo sapiens KIAA0431 mRNA partial cds	56	other
RC_AA416986	AA416986	Guanine nucleotide binding protein (G protein) beta polypeptide 1	56	TM
RC_AA435742_s	AA435742	Human fatty acid amide hydrolase mRNA complete cds	56	TM
RC_AA435936	AA435936	EST	56	7
RC_AA436819	AA436819	ESTs	56	other
RC_AA452842	AA452842	ESTs	56	other
RC_AA453987	AA453987	ESTs	56	other
AA477214	AA477214	ESTs	56	other
RC_AA482269	AA482269	Integral transmembrane protein 1	56	TM
D50914	D50914	Human mRNA for KIAA0124 gene partial cds	56	TM

FIGURE 7 (cont.)
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20 1151222		FOT.		
RC_N51590_s	N51590	ESTS	5.6	other
RC_N93797	N93797	ESTs SOT A Market No. 1974 And A Market No.	56	SS,
RC_T23528	T23528	ESTs Moderately similar to TYKi protein [M.musculus]	5.6	other
RC_T95057_f	T95057	ESTS	5.6	other
U91327	U91327	EST - U91327	5.6	?
X76105	X76105	H.sapiens DAP-1 mRNA	5.6	other
RC_AA021182	AA021182	ESTs	55	other
RC_AA075200	AA075200	Homo sapiens Chromosome 16 BAC clone CIT987SK-A-319E8	5.5	other
RC_AA085589	AA085589	ESTs Highly similar to TRANSLATION INITIATION FACTOR EIF-2B-DELTA SUBUNIT [Oryctolagus cuniculus]	55	other
RC_AA115535	AA115535	ESTs	5.5	other
RC_AA195517	AA195517	ESTs Weakly similar to !!!! ALU SUBFAMILY J WARNING ENTRY !!! [H.sapiens]	5.5	TM
RC_AA280840	AA280840	EŞTs	55	SS,
RC_AA443602	AA443602	ESTs	5.5	TM
RC_AA609996	AA609996	ESTs Highly similar to Surf-4 protein [M.musculus]	5 5	?
RC_H99500	H99500	Homo sapiens mRNA for follistain-related protein (FRP) complete cds	5.5	other
M24486	M24486	Procollagen-profine 2-oxoglutarate 4-dioxygenase (profine 4-hydroxylase) alpha	5.5	SS.
RC_N21032	N21032	polypeptide EST		
_	N22015	ESTs	55	?
RC_N22015		ESTs .	5.5	TM
RC_R43883	R43883		5.5	other
U09278	U09278	Human fibroblast activation protein mRNA complete cds	5.5	SS,
X02874	X02874	(2'-5') oligoadenylate synthetase E	5.5	other
RC_AA242757	AA242757	ESTs	5.4	other
RC_AA251973	AA251973	ESTs	5.4	?
RC_AA293773	AA293773	Homo sapiens clone 23870 mRNA sequence	5.4	other
RC_AA398721	AA398721	EST - RC_AA398721	5.4	other
RC_AA437225	AA437225	ESTs	54	other
RC_AA448410	AA448410	ESTs	5 4	other
RC_AA449357	AA449357	ESTs	5.4	other
RC_AA479348_s	AA479348	H sapiens mRNA for SYT	5 4	other
C02170	C02170	ESTs Weakly similar to weak similarity to ribosomal protein L14 [C elegans]	5 4	other
RC_H75933_i	H75933	Laminin receptor (2H5 epitope)	5 4	other
M97936	M97936	SIGNAL TRANSDUCER AND ACTIVATOR OF TRANSCRIPTION 1-ALPHA/BETA	5.4	other
RC_N51917	N51917	ESTs	5 4	other
RC_R41294_s	R41294	ESTs	5 4	other
RC_T81310	T81310	EST	5.4	?
RC_W45275_f	W45275	CD44 antigen (cell adhesion molecule)	5 4	TM
RC_W92001	W92001	ESTs	5 4	other
RC_AA135809	AA135809	ESTs	53	other
RC_AA157811	AA157811	EST - RC_AA157811	53	TM
RC_AA191524	AA191524	ESTs	53	other
RC_AA195036_s	AA195036	Human Ro/SSA ribonucleoprotein homolog (RoRet) mRNA complete cds	53	7
RC_AA284565_f	AA284565	ESTs	53	other
RC_AA287022_s	AA287022	Thymidine kinase 1 soluble	53	other
RC_AA394071	AA394071	Homo sapiens gamma2-adaptın (G2AD) mRNA complete cds	5.3	other
RC_AA399477	AA399477	ESTs	53	2
RC_AA401428_s	AA401428	NUCLEAR PORE COMPLEX PROTEIN NUP214	5.3	TM
RC_AA425154	AA425154	ESTs	5.3	other
RC_AA447213_s	AA447213	ESTs Weakly similar to 50S RIBOSOMAL PROTEIN L20 [E coli]	53	other
RC_AA464860	AA464860	Homo sapiens Jak2 kinase mRNA complete cds	53	other
RC_AA465191	AA465191	ESTs	5 3	other
RC_AA476293	AA476293	ESTs Weakly similar to DNA-DIRECTED RNA POLYMERASE III LARGEST SUBUNIT	53	TM
RC_H80749	H80749	[Plasmodium falciparum] ESTs	53	other
RC_N94385_s	N94385	Human germline oligomeric matrix protein (COMP) mRNA complete cds	53	other
U59877	U59877	Human low-Mr GTP-binding protein (RAB31) mRNA complete cds	53	other
X03363	X03363	ERBB-2 RECEPTOR PROTEIN-TYROSINE KINASE PRECURSOR	53	TM
RC_AA005262	AA005262	Homo sapiens DNA sequence from PAC 262D12 on chromosome q23 3-24 3 Contains	52 .	other
_		a Tenascin (Hexabrachion Cytotactın Neuronectin Myotendinous antigen)-LIKE gene and a mitochondrial/chloroplast 30S ribosom		
RC_AA025370	AA025370	ESTs	5 2	?
RC_AA026682_s	AA026682	Topoisomerase (DNA) II alpha (170kD)	5 2	other
RC_AA279160	AA279160	ESTs	52	7
RC_AA403116	AA403116	Homo sapiens U-snRNP-associated cyclophilin (USA-CyP) mRNA complete cds	52	other
RC_AA452857	AA452857	ESTs ,	5 2	?
RC_AA488280	AA488280	EST - RC_AA488280	5 2	other

FIGURE 7 (cont.)
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00.44500440		FOT. Market and the FINOSO PROTEIN THE		
RC_AA599140	AA599140 AA609891	ESTs Moderately similar to ZINC FINGER PROTEIN 7 [Homo sapiens] EST	5.2 5.2	other ?
RC_AA609891 RC_AA621714	AA621714	ESTs	5.2	other
H87319	H87319	Protein kinase C substrate 80K-H	5.2	TM
RC_H96392	H96392	ESTs	5.2	other
RC_N54321	N54321	EST	5,2	?
RC_N73861	N73861	EST - RC_N73861	5.2	other
RC_R05312_s	R05312	ESTs	5.2	other
RC_R59183_f	R59183	ESTs	5.2	other
RC_R92205	R92205	ESTs	5.2	TM
RC_W45302	W45302	ESTs Highly similar to HYPOTHETICAL HELICASE K12H4.8 IN CHROMOSOME III	5.2	other
X17644	X17644	[Caenorhabditis elegans] G1 to S phase transition 1	5.2	other
RC_AA284518	AA284518	ESTs	5.1	other
RC_AA434152	AA434152	ESTs	5,1	SS,
RC_AA436673	AA436673	ESTs	5.1	other
RC_AA481453	AA481453	ESTs	5.1	other
RC_AA487449	AA487449	EST - RC_AA487449	5,1	other
RC_AA491465	AA491465	ESTs	5.1	other
RC_N79612	N79612	ESTs	5.1	other
RC_N98461	N98461	ESTs	5.1	TM
RC_R42036	R42036	ESTs	5.1	other
RC_R43543	R43543	ESTs	5.1	other
RC_W59961_s	W59961	Human mRNA for KIAA0389 gene complete cds	5.1	other
RC_W60180	W60180	ESTs	5.1	other
Y09912_ma1	Y09912	H sapiens mRNA for AP-2 beta transcription factor	5.1	?
RC_AA055892	AA055892	ESTs	5	other
RC_AA085676	AA085676	ESTs Weakly similar to TYL [H.sapiens]	5	other
RC_AA172056	AA172056	ESTs	5	other
RC_AA211400	AA211400	ESTs	5	other
RC_AA236356	AA236356	ES7s	5	other
RC_AA252395	AA252395	ESTs	5	2
RC_AA262179	AA262179	ESTs	5	other
RC_AA281451	AA281451	ESTs	5	other
RC_AA287095	AA287095	EST - RC_AA287095	5	?
RC_AA425691	AA425691	ESTs	5	other
RC_AA426376	AA426376	ESTs	5	other
RC_AA446000	AA446000	ESTs	5	other
RC_AA478951	AA478951	EST	5	2
AA479995	AA479995	Homo sapiens mRNA for KIAA0583 protein partial cds	5	other
D82419	D82419	ESTs Highly similar to UBIQUITIN-CONJUGATING ENZYME E2-28 4 KD [Saccharomyces cerevisiae]	5	TM
RC_F02080_f	F02080	ESTs	5	other
RC_F10496_f	F10496	H.sapiens 40 kDa protein kinase related to rat ERK2	5	other
J00314	J00314	Homo sapiens clone 24703 beta-tubulin mRNA complete cds	5	?
J05614	J05614	EST - J05614	5	2
M16336	M16336	CD2 antigen (T cell surface antigen T11)	5	TM
M80244	M80244	INTEGRAL MEMBRANE PROTEIN E16	5	TM
RC_N33927_s	N33927	"Homo sapiens mRNA for histone H2B clone pJG4-5-15""	5	SS,
RC_N73808_f	N73808	ESTs	5	?
RC_T79815 U62392	T79815	ESTs Weakly similar to !!!! ALU SUBFAMILY J WARNING ENTRY !!!! [H sapiens]	5	other
RC_AA133756	U62392	Homo sapiens zinc finger protein mRNA complete cds	5	other
RC_AA234559	AA133756 AA234559	ESTs	49	other
RC_AA490830	AA490830	ESTs	49 49	other
C01169	C01169	Homo sapiens clone 23915 mRNA sequence	49	other ather
D21255	D21255	Cadherin 11 (OB-cadherin)	49	SS,TM
RC_F10945	F10945	Polypyrimidine tract binding protein (hnRNP I) (alternative products)	49	other
RC_H24044	H24044	Protein phosphatase 2 (formerly 2A) catalytic subunit alpha isoform	49	other
RC_N34893	N34893	ESTs Highly similar to HYPOTHETICAL 47.8 KD PROTEIN 80280.9 IN	49	other
RC_R41772		CHROMOSOME III (Caenorhabditis elegans)		
_	R41772	EST DO TRACE	49	?
RC_T59338	T59338	EST - RC_T59338	4.9	other
RC_AA191512	AA191512	ESTs EST-	48	?
RC_AA400513_i RC_AA406081	AA400513 AA406081	ESTs	48	other
	~~~0000 i	ESTs	48	other

FIGURE 7 (cont.)
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00 4449169	1149450	ECT	4.8	?
RC_AA448158 RC_N94362	AA448158 N94362	EST EST	48	?
RC_W60007_s	W60007	Human mRNA for KIAA0203 gene complete cds	48	other
RC_AA026356	AA026356	ESTs	47	7
AA075599	AA075599	ESTS Highly similar to NADH-UBIQUINONE OXIDOREDUCTASE B22 SUBUNIT (Bos taurus)	47	other
RC_AA157836	AA157836	ESTs	47	other
RC_AA196549.	AA196549	ESTs	47	other
RC_AA404352	AA404352	ESTs	4.7	other
RC_AA417321	AA417321	ESTs Weakly similar to CALMODULIN [D.melanogaster]	47	SS,
RC_AA418074	AA418074	ESTs	4.7	other
RC_N32919	N32919	ESTs	4.7	other
RC_AA177051	AA177051	EST - RC_AA177051	4.6	7
RC_AA453483	AA453483	ESTs	4.6	TM
RC_AA620795	AA520795	ESTs	46	SS,TM
RC_H97012	H97012	ESTs Weakly similar to LB004.7 gene product [S.cerevisiae]	46	other
M34458_ma1	M34458	LAMIN B1	4.6	other
RC_N68921	N68921	ESTs	4.6	other
U18321	U18321	H.sapiens DAP-3 mRNA	4.6	other
X17059	X17059	ARYLAMINE N-ACETYLTRANSFERASE MONOMORPHIC	4.6	?
RC_AA210722	AA210722	EST	45	?
RC_AA255605	AA255605	Homo sapiens spindle pole body protein spc97 homolog GCP2 mRNA complete cds	45	other
RC_AA443634	AA443634	Homo sapiens ubiquitin conjugating enzyme G2 (UBE2G2) mRNA complete cds	4.5	?
RC_AA461507	AA461507	ESTs .	4.5	TM
RC_AA464853	AA464853	ESTs Weakly similar to T01G9 4 [C.elegans]	4.5	other
RC_N71076	N71076	EST	45	?
RC_T40841	T40841	ESTs	4.5	other
X57766	X57766	Human stromelysin-3 mRNA	4.5	other
RC_AA206497_s	AA206497	PROTEASOME COMPONENT C9	4.4	other
RC_AA227900_s	AA227900	H.sapiens mRNA homologous to S, cerevisiae RAD54	4.4	other
RC_AA346495	AA346495	ESTs Moderately similar to !!!! ALU SUBFAMILY J WARNING ENTRY !!!! [H.sapiens]	4,4	?
RC_AA386260	AA386260	EST	4.4	?
RC_AA398155	AA398155	ESTs	4.4	other
RC_AA405569_s	AA405569	Human fibroblast activation protein mRNA complete cds	4 4	SS.
AA422025_s	AA422025	ESTs	4 4	other
RC_AA430124	AA430124	ESTs	4 4	other
RC_AA453466	AA453466	ESTs	4 4	other
RC_AA463726_s	AA463726	Homo sapiens mRNA for JM27 protein complete CDS (clone IMAGE 145745 and IMAGE 257878)	4.4	other
RC_C20981	C20981	ESTs Highly similar to CHOLINE DEHYDROGENASE [Escherichia coli]	44	other
RC_R70801_s	R70801	EST DO TOTOOZ	4.4	other
RC_T97307 U28386	T97307 U28386	EST - RC_T97307	4 4 4 4	7
X02419_ma1	X02419	RAG (recombination activating gene) cohort 1 Urokinase-type plasminogen activator	44	TM ?
RC_AA235112	AA235112	ESTs	43	r other
RC_AA453176_s	AA453176	Human protein kinase ATR mRNA complete cds	4.3	TM
D42073	D42073	Human mRNA for reticulocalbin complete cds	43	SS,
U21090	U21090	Human DNA polymerase delta small subunit mRNA complete cds	43	other
U24389	U24389	Human lysyl oxidase-like protein mRNA complete cds	4,3	7
U73379	U73379	Human cyclin-selective ubiquitin carrier protein mRNA complete cds	43	other
RC_AA227959_s	AA227959	Human cysteine protease Mch2 isoform alpha (Mch2) mRNA complete cds	4.2	other
RC_AA416931	AA416931	ESTs	42	other
RC_AA419200	AA419200	ESTs	42	SS.
RC_H18947	H18947	ESTs	42	other
RC_H25577	H25577	ESTs Highly similar to CYTOCHROME P450 IVB1 [Oryctolagus cuniculus]	42	TM
RC_H90161_s	H90161	ESTs	4.2	SS,
M15796	M15796	Proliferating cell nuclear antigen	42	?
RC_R46482	R46482	ESTs	42	other
S74445	S74445	Cellular retinoic acid-binding protein [human skin mRNA 735 nt]	42	other
U74612	U74612	Human putative M phase phosphoprotein 2 (MPP2) mRNA complete cds	42	TM
X52534	X62534	High-mobility group (nonhistone chromosomal) protein 2	42	other
RC_AA398369	AA398369	ESTs	41	other
RC_AA448347	AA448347	Annexin XI (56kD autoantigen)	4 1	other
RC_AA464707	AA464707	ESTs	41	TM
RC_AA478799_s	AA478799	H sapiens mRNA for BS69 protein	41	TM

FIGURE 7 (cont.)
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RC_AA496369	AA496369	ESTs	41	other
RC_D54296_f	D54296	Human mRNA for KIAA0255 gene complete cds	4.1	TM
RC_N66818 U50648	N66818 U50648	ESTs Protein kinase interferon-inducible double stranded RNA dependent	4.1	TM ?
AA193297	AA193297	ESTs	4 1 4	SS,
RC_AA228026	AA228026	ESTs Highly similar to PBDX protein [H.sapiens]	4	TM
RC_AA287325_f	AA287325	ESTs	4	?
RC_AA287596	AA287596	ESTs	4	t other
RC_AA421041	AA421041	ESTs	4	other
U29463	U29463	Cytochrome B561	4	?
RC_W87752_s	W87752	Small inducible cytokine A5 (RANTES)	4	TM
X94563_xpt2_r	X94563	EST - X94563_xpt2_r	4	?
RC_AA256837_i	AA256837	ESTs	3.9	?
RC_AA416627_s	AA416627	ESTs	3.9	other
RC_AA482224_f	AA482224	ESTs Weakly similar to No definition line found [C.elegans]	3.9	7
RC_AA485360	AA485360	EST	39	?
RC_R44709	R44709	Homo sapiens mRNA for RB18A protein	3.9	other
RC_W45572_f	W4\$572	ADP-ribosylation factor 1	3.9	other
RC_AA132366	AA132366	Homo sapiens mRNA for SPOP	3.8	?
RC_AA133527	AA133527	ESTs Moderately similar to The KIAA0138 gene product is novel. [H.sapiens]	3.8	other
RC_AA224324	AA224324	ESTs	38	other
RC_AA287642_s	AA287642	Human mRNA for KIAA0078 gene complete cds	38	other
RC_AA425652	AA425652	ESTs	3.8	other
RC_AA459960_s	AA459960	ESTs Weakly similar to D9481.16 gene product [S.cerevisiae]	3.8	other
RC_AA465094	AA465094	ESTs Weakly similar to nemo form II [D.melanogaster]	3.8	other
RC_AA485451	AA485451	EST	38	other
RC_AA599244	AA599244	Homo sapiens mRNA for KIAA0530 protein partial cds	3.8	other
RC_N41018	N41018	Human mRNA for prepro cortistatin like peptide complete cds	38	?
RC_N74501	N74501	ESTs	38	other
RC_N98525	N98525	Homo sapiens tumorous imaginal discs protein Tid56 homolog (TID1) mRNA complete cds	3.8	other
W46488	W46488	Homo sapiens Amplified in Breast Cancer (AIB1) mRNA complete cds	38	other
RC_AA232183	AA232183	ESTs Weakly similar to IIII ALU SUBFAMILY J WARNING ENTRY IIII [H sapiens]	37	other
RC_AA399547	AA399547	ESTs	37	other
RC_AA424486	AA424486	ESTs	37	TM
RC_AA598661	AA598661	ESTs	37	other
RC_H04339	H04339	ESTs	37	TM
M14219	M14219	Decorin	3.7	other
M25077	M25077	Human 60-kdal ribonucleoprotein (Ro) mRNA complete cds	3.7	?
RC_N67102_s	N67102	ESTs	37	other
RC_W45728	W45728	ESTS Highly similar to HETEROGENEOUS NUCLEAR RIBONUCLEOPROTEIN L	37	other
Y12394	Y12394	[Homo sapiens] Homo sapiens importin-alpha homolog (SRP1gamma) mRNA complete cds	37	other
RC_AA401758_i	AA401758	ESTs Weakly similar to !!!! ALU SUBFAMILY SQ WARNING ENTRY !!!! [H sapiens]	36	SS,
RC_AA435840	AA435840	Homo sapiens mRNA for high mobility group protein HMG2a	36	other
RC_AA461492	AA461492	ESTs	36	other
RC_AA521240	AA521240	ESTs	36	TM
RC_F02450	F02450	ESTs Moderately similar to unknown protein [H sapiens]	36	TM
RC_N63823	N63823	ESTs	36	other
RC_N67603	N67603	ESTs Weakly similar to hypothetical L1 protein [H sapiens]	36	7
RC_N91687_s	N91887	Homo sapiens mRNA for NB thymosin beta complete cds	36	other
RC_N93967	N93967	EST	36	other
RC_W73788	W73788	ESTs	36	other
RC_AA034365	AA034365	NUCLEAR PORE GLYCOPROTEIN P62	35	other
RC_AA083069	AA083069	EST - RC_AA083069	3 5	7
RC_AA112063	AA112063	ESTs Weakly similar to PRE-MRNA SPLICING HELICASE BRR2 [S.cerevisiae]	35	other
RC_AA126951	AA126951·	ESTs Weakly similar to DNA-directed RNA polymerase [D melanogaster]	35	other
RC_AA159181	AA159181	ESTs Weakly similar to Lpa8p [S cerevisiae]	35	other
RC_AA398450	AA398450	H sapiens mRNA for synaptonemal complex lateral element protein	35	other
RC_AA404593	AA404593	ESTs	35	other
RC_AA412739	AA412739	EST	35	other
RC_AA447626	AA447526 AA453787	EST  Number TEIIB soleled forder hBBC (UBBC) - BNA complete and	35	?
RC_AA453787_s RC_AA599106	AA453787 AA599106	Human TFIIB related factor hBRF (HBRF) mRNA complete cds EST - RC_AA599106	35	other
D82558	D82558	_	35	other
_ >2.000	202000	Homo sapiens KB07 protein mRNA partial cds	3 5	other

FIGURE 7 (cont.)
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RC_H72283_s	H72283	Human mRNA for KIAA0265 gene partial cds	3.5	other
L38961	L38961 N36835	Integral transmembrane protein 1 ESTs	35 35	TM other
RC_N36835 RC_N90859	N90859	ESTs	35	other
RC_R63734	R63734	ESTs	35	TM
R70167	R70167	ESTs	35	other
X69141	X69141	FARNESYL-DIPHOSPHATE FARNESYLTRANSFERASE	3.5	other
X75346	X75346	H.sapiens mRNA for MAP kinase activated protein kinase	35	TM
RC_AA029042	AA029042	Human hStAH2 mRNA complete cds	3.4	other
RC_AA100470	AA100470	ESTs	3.4	other
AA115397	AA115397	Homo sapiens mRNA for putative methyltransferase	3.4	other
RC_AA164209	AA164209	Homo sapiens RRM RNA binding protein Gry-rbp (GRY-RBP) mRNA complete cds	34	other
RC_AA258203	AA258203	ESTs	3.4	other
RC_AA411448_s	AA411448	ESTs	3.4	TM
RC_AA429917	AA429917	ESTs	3.4	TM
RC_AA442070_s	AA442070	Phosphoribosyl pyrophosphate amidotransferase	3.4	other
RC_AA449417	AA449417	Homo sapiens mRNA for putative glucosyltransferase partial cds	3.4	TM
RC_AA453164	AA453164	EST	3.4	?
RC_F10326_f	F10326	EST	3.4	other
RC_H88639	H88639	ESTs	3.4	other
L47276	L47276	EST - L47276	3.4	other
RC_N29740	N29740	ESTs	3.4	other
RC_N33920	N33920	H.sapiens mRNA for diubiquitin	3.4	other
RC_N34895	N34895	ESTs	3.4	other
S81003	\$81003	L-UBC	3.4	other
U07806	U07806	DNA topoisomerase I	3 4	other
RC_AA047896	AA047896	ESTS ESTS	33	other
RC_AA116036	AA116036 AA232535		33	other
RC_AA232535_s	AA202000	ESTs Weakly similar to LINE-1 REVERSE TRANSCRIPTASE HOMOLOG [Homo sapiens]	33	other
RC_AA435847	AA435847	EST - RC_AA435847	33	other
RC_AA453159_s	AA453159	Human kinesin-like spindle protein HKSP (HKSP) mRNA complete cds	3.3	other
RC_AA490899	AA490899	ESTs	33	other
RC_AA496051	AA496051	ESTs	33	TM
AD000092_cds7 RC_F09353	AD000092 F09353	Homo sapiens DNA from chromosome 19p13 2 cosmids R31240 R30272 and R28549 containing the EKLF GCDH CRTC and RAD23A genes genomic sequence Homo sapiens sodium/myo-inositol cotransporter (SLC5A3) gene complete cds	3 3 3 3	? other
RC_N34059	N34059	EST - RC_N34059	33	other
RC_N58172	N58172	ESTs Weakly similar to IIII ALU SUBFAMILY SC WARNING ENTRY IIII [H sapiens]	33	TM
RC_N67437	N67437	ESTs	33	TM
RC_R24237_f	R24237	ESTs	33	?
RC_R45356	R45356	Homo sapiens cDNA similar to RNA binding protein C elegans complete	33	other
RC_W44735	W44735	ESTs	33	2
RC_W85861	W85861	ESTs Weakly similar to ZK1058.4 [C elegans]	33	SS,TM
RC_AA134965_i	AA134965	ESTs	3 2	other
RC_AA169379	AA169379	ESTs	32	other
RC_AA211941 RC_AA232939	AA211941 AA232939	Homo sapiens polyadenylate binding protein-interacting protein-1 (PAIP1) mRNA complete cds ESTs	3 2 3 2	other
AA421213	AA421213	ESTs Weakly similar to F28F8 3 [C elegans]		other other
RC_AA422079	AA422079	ESTs Weakly similar to RAR-RESPONSIVE PROTEIN TIG1 [H sapiens]	3 2 3 2	other
RC_AA448213_s	AA448213	Human myogenic repressor I-mf (MDFI) mRNA complete cds	32	TM
RC_AA490969	AA490969	ESTs	32	other
RC_AA609423	AA609423	ESTs	32	other
D84145	D84145	Human WS-3 mRNA complete cds	32	other
RC_F09315	F09315	Homo sapiens mRNA for KIAA0583 protein partial cds	32	other
L07515	L07515	HETEROCHROMATIN PROTEIN 1 HOMOLOG	32	other
M86852	M86852	Peroxisomal membrane protein 3 (35kD Zeilweger syndrome)	32	TM
RC_N35385	N35385	ESTs	32	other
RC_N78572	N78572	EST - RC_N78572	32	other
RC_R60192_s	R60192	Peroxisomal biogenesis factor 7	32	other
RC_R67996	R67996	ESTs .	32	other
RC_T15665	T15665	ESTs Weakly similar to HYPOTHETICAL 139 1 KD PROTEIN CO8B11 3 IN CHROMOSOME II [C elegans]	32	other
T40327_s	T40327	ESTs	32	other
RC_T65797	T65797	ESTs Weakly similar to Pin1 protein [H sapiens]	32	TM

FIGURE 7 (cont.)
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U37547	U37547	Himan IAD hamalag B (MINIO) - DNA samplete ada	2.2	-41
U51586	U51586	Human IAP homolog B (MIHB) mRNA complete cds	3 2	other
U81554	U81554	Human siah binding protein 1 (SiahBP1) mRNA partial cds Homo sapiens signal recognition particle 72 (SRP72) mRNA complete cds	3.2	other
RC_W33134_s	W33134	ESTs	3.2 3.2	other ?
RC_W46255	W46255	ESTs	3.2	other
Z97054_xpt2	Z97054	Human mRNA for KIAA0312 gene partial cds	3,2	?
RC_AA047036	AA047036	ESTs	3.1	olher
RC_AA150043	AA150043	ESTs	3.1	TM
RC_AA232874	AA232874	EST	31	TM
RC_AA291259	AA291259	ESTs	3.1	TM
RC_AA398360	AA398360	EST	3.1	7
RC_AA401687_s	AA401687	Homo sapiens ribonuclease P protein subunit p20 (RPP20) mRNA complete cds	31	other
RC_D60208_f	D60208	ESTs	3.1	other
D87466	D87466	Human mRNA for KtAA0276 gene partial cds	3.1	other
RC_N21626	N21626	ESTs	3,1	other
RC_N69331	N69331	Peptidylprolyl isomerase C (cyclophilin C)	3.1	TM
\$66431	S66431	Homo sapiens clone 23592 mRNA sequence	3.1	other
RC_W44928	W44928	ESTs	31	TM
RC_W72967	W72967	ESTs	3.1	other
X17620	X17620	NUCLEOSIDE DIPHOSPHATE KINASE A	3.1	other
X59798	X59798	Cyclin D1 (PRAD1 parathyroid adenomatosis 1)	31	other
RC_AA099719	AA099719	EST\$	3	other
RC_AA152305_s	AA152305	Interferon (gamma)-induced cell line protein 10 from	3	SS,
RC_AA227932	AA227932	ESTs	3	other
RC_AA251738	AA251738	H.sapiens mRNA for TAFII100 protein	3	other
RC_AA386264	AA386264	ESTs Highly similar to ribosome-binding protein p34 [R norvegicus]	3	other ,
RC_AA406577	AA406577	ESTs	3	other
RC_AA423827_r	AA423827	ESTs	3	other
RC_N47204	N47204	ESTs Weakly similar to C50F4 12 [C.elegans]	3	other
RC_R36548	R36548	ESTs	3	TM
\$50223	\$50223	HKR-T1	3	other
RC_W46286_s	W46286	ESTs Weakly similar to ZK1058.5 [C elegans]	3	TM
RC_W80482	W80482	ESTs	3	other
X70944_s	X70944	PTB-ASSOCIATED SPLICING FACTOR	3	other
X74801	X74801	H sapiens Cdg mRNA for chaperonin	3	other
Y12065	Y12065	Homo sapiens mRNA for nucleolar protein hNop56	3	?
RC_AA164293_f	AA164293	ESTs	29	?
RC_AA179845	AA179845	ESTs Moderately similar to rabkinesin-6 [M musculus]	29	other
RC_AA195936	AA195936	ESTs Weakly similar to W02D9 2 [C elegans]	29	TM
AA203523	AA203523	ESTs Weakly similar to coded for by C elegans cDNA yk10c10 3 [C elegans]	29	other
RC_AA206088	AA206088	ESTs	29	other
RC_AA213506	AA213506	ESTs	29	other
RC_AA228020	AA228020	Homo sapiens splicing factor (CC1 3) mRNA complete cds	2.9	other
RC_AA242834	AA242834	ESTs	29	other
RC_AA279420	AA279420	ESTs Weakly similar to T08A11.2 [C elegans]	29	TM
RC_AA292747	AA292747	ESTs	29	other
AA393164_s	AA393164	Homo sapiens mammaglobin B precursor mRNA complete cds	2.9	TM
RC_AA399164	AA399164	ESTs	29	other
RC_AA399264	AA399264	ESTs	29	other
RC_AA400725	AA400725	ESTs	29	other
RC_AA426120	AA426120	EST - RC_AA426120	29	other
AA452011 RC_AA485214_s	AA452011 AA485214	ESTS Highly similar to deduced protein product shows significant homology to coactosin from Dictyostellum discoideum [H sapiens] DNA-BINDING PROTEIN NEFA PRECURSOR	29 29	other
D00596	D00596	Thymidylate synthase	2.9	SS, ?
RC_D60061_s	D60061	ESTs		
RC_F13779	F13779	ESTs	2.9 2.9	other
RC_H16790	H16790	ESTs	2.9 2.9	SS,
RC_H97677_s	H97677	ESTs	29	other other
M28211	M28211	Homo sapiens GTP-binding protein (RAB4) mRNA complete cds	2.9	other
RC_N32333	N32333	Homo sapiens M962 protein spliced isoform 2 mRNA complete cds	2.9	other
RC_N36881	N36881	ESTs	29	other
RC_R50840	R50840	H sapiens mRNA for ras-related GTP-binding protein	29	other
RC_R97040	R97040	ESTs	29	other
_				0

FIGURE 7 (cont.)
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RC_T25732_f	T25732	Human mRNA for KIAA0252 gene partial cds	29	?
T54762_s	T54762	ESTs	2.9	?
HG110-HT110	TIGR - HG110- HT110	EST - HG110-HT110	2.9	?
U40714	U40714	Human tyrosyl-IRNA synthetase mRNA complete cds	2.9	other
RC_AA001409_i	AA001409	ESTs	2.8	other
RC_AA128407	AA128407	ESTs	28	other
RC_AA232231	AA232231	ESTs	28	other
RC_AA262768	AA262768	ESTs	2.8	TM
RC_AA292765	AA292765	H.sapiens mRNA for M-phase phosphoprotein mpp5	28	other
RC_AA310729_s	AA310729	Human mRNA for clathrin-like protein complete cds	28	TM
RC AA405512	AA405512	ESTs	2.8	other
RC_AA411532	AA411532	ESTs Weakly similar to ORF YOR285w [S.cerevisiae]	2.8	other
_		EST	2.8	?
RC_AA412497	AA412497			
RC_AA425606	AA425606	ESTs Weakly similar to Similar to S.cerevisiae hypothetical protein L3111 [H sapiens]	2.8	TM
RC_AA425900_s	AA425900	Uracil-DNA glycosylase	28	other
RC_AA446572	AA446572	EST - RC_AA446572	2.8	other
RC_AA478596	AA478596	ESTs	28	?
RC_AA480103	AA480103	ESTs Weakly similar to !!!! ALU SUBFAMILY J WARNING ENTRY !!!! [H.sapiens]	2.8	TM
RC_AA486407	AA486407	ESTs	2.8	other
RC_AA488432	AA488432	ESTs	2.8	7
RC_AA609200	AA609200	EST - RC_AA609200	2.8	other
RC_AA609501	AA609501	HEAT SHOCK 70 KD PROTEIN 1	28	other
AF006516	AF006516	Homo sapiens eps8 binding protein e3B1 mRNA complete cds	28	other
RC_D80710_f	D80710	ESTs Weakly similar to transmembrane protein [H.sapiens]	2.8	? `
RC_F03605_f	F03605	PUTATIVE 60S RIBOSOMAL PROTEIN	2.8	other
RC_H24460_s	H24460	FK506-binding protein 4 (59kD)	28	other
RC_N21159	N21159	Homo sapiens forkhead protein (FKHRL1) mRNA complete cds	2.8	other
RC_N29325	N29325	ESTs Highly similar to 47 KD PROTEIN [Pseudomonas chlororaphis]	28	other
RC_N48715	N48715	ESTs	28	SS,TM
RC_N92915	N92915	ESTs	2.8	other
RC_R39234_r	R39234	ESTs Weakly similar to elastin like protein [D melanogaster]	2.8	TM
RC_R41933	R41933	ESTs	2.8	2
RC_R46025	R46025	ESTs	2.8	SS.
RC_R49327	R49327	Natural resistance-associated macrophage protein 2	28	TM
RC_R79617	R79617	ESTs	28	other
<del>-</del>			28	?
RC_T63857	T63857	EST - RC_T63857		
U30825	U30825	Human splicing factor SRp30c mRNA complete cds	28	other
U53347	U53347	Human neutral amino acid transporter B mRNA complete cds	28	TM
X76732	X76732	DNA-BINDING PROTEIN NEFA PRECURSOR	28	SS,
RC_AA102520 RC_AA125969	AA102520 AA125969	ESTs Highly similar to HYPOTHETICAL 31.6 KD PROTEIN F54F2 9 IN CHROMOSOME III [Caenorhabditis elegans] ESTs Weakly similar to F35G12.9 [C elegans]	27 27	TM 2
-			27	
RC_AA126743	AA126743	ESTs		other
RC_AA164687	AA164687	ESTs	27	other
AA215333	AA215333	ESTs	27	TM.
RC_AA291269	AA291269	ESTs	2.7	other
RC_AA424031	AA424031	ESTs	27	other
RC_AA425725	AA425725	ESTs Weakly similar to serine protein kinase SRPK1 [H sapiens]	27	other
RC_AA431333_s	AA431333	ESTs Highly similar to Ras inhibitor [H sapiens]	27	other
RC_AA449718	AA449718	ESTs Weakly similar to ZINC FINGER PROTEIN 42 [H sapiens]	27	other
RC_AA461509	AA461509	ESTs Weakly similar to putative p150 [H sapiens]	27	other
RC_AA620586	AA620586	ESTs	27	2
D63391	D63391	Human mRNA for platelet activating factor acetylhydrolase IB gamma-subunit complete cds	27	other
RC_F02863	F02863	ESTs Moderately similar to !!!! ALU SUBFAMILY SQ WARNING ENTRY !!!! [H sapiens]	27	TM
K02777	K02777	T cell receptor alpha-chain	27	MT
L11669	L11669	Human tetracycline transporter-like protein mRNA complete cds	27	SS,TM
L12350	L12350	Thrombospondin 2	27	other
M96982	M96982	SPLICING FACTOR U2AF 35 KD SUBUNIT	27	7
RC_N23663	N23663	ESTs	27	other
RC_N25798	N25798	ESTs	2.7	ТМ
RC_R02354	R02354	ESTs	27	other
RC_R54112	R54112	ESTs	27	other
RC_R71481	R71481	ESTs	27	TM
U33052	U33052	Human lipid-ectivated protein kinase PRK2 mRNA complete cds	27	other

FIGURE 7 (cont.)
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RC_W69160	W69160	ESTs	27	other
RC_W80750	W80750	ESTs	27	other
RC_W87747	W87747	ESTs	2.7	other
X16396	X16396	NAD-DEPENDENT METHYLENETETRAHYDROFOLATE DEHYDROGENASE	27	SS.
X89986	X89986	H.sapiens mRNA for NBK apoptotic inducer protein	27	TM
RC_AA001402	AA001402	Homo sapiens 15 kDa selenoprotein mRNA complete cds	26	other
RC_AA176121	AA176121	ESTs	26	other
RC_AA242758_s	AA242758	Human breast cancer estrogen regulated LIV-1 protein (LIV-1) mRNA partial cds	2.6	SS,TM
RC_AA293300_s	AA293300	ESTs Weakly similar to semaphorin C [M.musculus]	2.6	SS.
RC_AA412112	AA412112	EST - RC_AA412112	2.6	?
RC_AA417956	AA417956	ESTs	2.6	other
RC_AA447553	AA447553	ESTs .	2.6	other
RC_AA447617	AA447617	ESTs	26	other
RC_AA453624	AA453624	Human terminal transferase mRNA complete cds	2.6	other
RC_AA598648_s	AA598648	Human mRNA for transcriptional activator hSNF2b complete cds	2.6	other
AB000115	AB000115	Homo sapiens mRNA expressed in osteoblast complete cds	2.6	other
AF001294	AF001294	Homo sapiens IPL (IPL) mRNA complete cds	2.6	other
D63881	D63881	Human mRNA for KIAA0160 gene partial cds	2.6	SS.
D79997	D79997	Human mRNA for KIAA0175 gene complete cds		
RC_H98655	H98655	Homo sapiens nibrin (NBS) mRNA complete cds	2.6	TM
L03411	L03411	Radin blood group	2.6 2.6	TM
		Homo sapiens (clone CC6) NADH-ubiquinone oxidoreductase subunit mRNA 3' end cds		other
L04490 BC N33546	L04490 N33516		2.6	other
RC_N33516		Homo sapiens nibrin (NBS) mRNA complete cds ESTs	26	TM
RC_N46252	N46252		2.6	TM
RC_N48790	N48790	ESTs Moderately similar to !!!! ALU SUBFAMILY SX WARNING ENTRY !!!! [H sapiens]	26	other
RC_N51316	N51316	ESTs Highly similar to elastin like protein [D.melanogaster]	26	other
RC_N63210	N63210	ESTs	26	other
RC_N68869	N68869	ESTS	2.6	other
RC_N92948_s	N92948	Human IEF SSP 9502 mRNA complete cds	26	other
RC_R50333_i	R50333	ESTs	26	?
T39176_s	T39176	ESTs Weakly similar to ZK1058.4 [C.elegans]	26	SS,TM
RC_W85712	W85712	ESTs Weakly similar to PROCOLLAGEN ALPHA 2(IV) CHAIN PRECURSOR [H.sapiens]	26	TM
Z49099	Z49099	H sapiens mRNA for spermine synthase	2.6	other
RC_AA045083	AA045083	VITAMIN K-DEPENDENT GAMMA-CARBOXYLASE	25	other
RC_AA069547	AA069547	EST - RC_AA069547	25	other
RC_AA130349	AA130349	ESTs	25	7
RC_AA160890_s	AA160890	Human mRNA for KiAA0389 gene complete cds	25	other
RC_AA191424	AA191424	ESTs	2.5	other
RC_AA236489	AA236489	ESTs	25	other
RC_AA251587	AA251587	Homo sapiens mRNA for KIAA0530 protein partial cds	25	other
RC_AA262491	AA262491	ESTs	25	other
RC_AA262730	AA262730	ESTs	25	?
RC_AA284372	AA284372	ESTs	25	other
RC_AA291503	AA291503	EST	2.5	?
RC_AA369027	AA369027	ESTs	25	SS,
RC_AA398280	AA398280	ESTs	2 5	other
RC_AA404957	AA404957	Matrix Gla protein	25	other
RC_AA416877	AA416877	ESTs	25	other
RC_AA428179	AA428179	EST	25	?
RC_AA446100	AA446100	ESTs	25	other
RC_AA451707	AA451707	ESTs	25	other
RC_AA490882_s	AA490882	ESTs	25	other
RC_AA610073	AA610073	ESTs	25	other
AB002308	AB002308	Human mRNA for KIAA0310 gene complete cds	25	other
D43948	D43948	Human mRNA for KIAA0097 gene complete cds	2.5	TM
RC_D60374_1	D60374	EST - RC_D60374_f	25	other
RC_H12634	H12634	ESTs	25	other
L00205	L00205	KERATIN TYPE II CYTOSKELETAL 6D	25	7
M23379	M23379	GTPase-activating protein ras p21 (RASA)	25	TM
RC_N21677	N21677	ESTs	25	other
RC_N66158	N66158	ESTs	2.5	TM
RC_N67187_s	N67187	ESTs	2.5	other
RC_N70646	N70646	ESTs	25	TM

FIGURE 7 (cont.)
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RC_N93000	N93000	ESTs	2.5	other
T39763_s	T39763	ESTs	25	other
RC_T70541	T70541	ESTs	2.5	SS,
U16306	U16306	LARGE FIBROBLAST PROTEOGLYCAN PRECURSOR	2.5	SS.
RC_W47183	W47183	ESTs	2.5	other
X54941	X54941	CDC28 protein kinase 1	2.5	other
X70218	X70218	Protein phosphatase 4 (formerly X) catalytic subunit	25	other
X85373	X85373	H sapiens mRNA for Sm protein G	25	other
RC_AA039887	AA039887	ESTs	2.4	other
RC_AA147708	AA147708	ESTs Highly similar to VACUOLAR ATP SYNTHASE SUBUNIT D [Bos taurus]	2.4	other
RC_AA190993	AA190993	ESTs ,	2.4	other
RC_AA218663	AA218663	ESTs Weakly similar to U1 SMALL NUCLEAR RIBONUCLEOPROTEIN 70 KD (Xenopus laevis)  EST Meakly similar to D0491 15 copp gradual IS coppidities)	2.4	other
RC_AA223209	AA223209	ESTs Weakly similar to D9481.16 gene product [S.cerevisiae]	24	other
RC_AA252672_s	AA252672	Homo sapiens diphthamide biosynthesis protein-2 (DPH2) mRNA complete cds	2.4	other
RC_AA258601	AA258601	EST - RC_AA258601	2.4	?
RC_AA262651	AA262651	ESTs	2.4	other
RC_AA279799	AA279799	ESTs	24	other
RC_AA286942	AA286942	EST - RC_AA286942	2.4	?
RC_AA371604	AA371604	Human Rho-associated coiled-coil containing protein kinase p160ROCK mRNA complete cds	2.4	other
RC_AA399047	AA399047	ESTs	2.4	other
AA434329	AA434329	ESTs	2.4	other
RC_AA449458	AA449458	ESTs	24	TM
RC_AA455239	AA455239	ESTs Highly similar to CHROMOSOME CONDENSATION PROTEIN DPY-27 [Caenorhabditis elegans]	24	other
RC_AA456646	AA456646	ESTs	24	other
RC_AA487207	AA487207	EST - RC_AA487207	2.4	other
AA504223 RC_AA599674	AA504223 AA599674	ESTs Highly similar to CHROMOSOME CONDENSATION PROTEIN DPY-27 [Caenorhabditis elegans] ESTs Weakly similar to F08G12.1 [C.elegans]	2 4 2 4	other SS.
-				
D38555	D38555	Human mRNA for KIAA0079 gene complete cds	24	other
D82348 D87684	D82348 D87684	Human mRNA for 5-aminoimidazole-4-carboxamide-1-beta-D-nbonucleot: de transformylase/inosinicase complete cds Human mRNA for KIAA0242 gene partial cds	2 4 2 4	other other
M55542	M55542			
M90516	M90516	Guanylate binding protein 1 interferon-inducible 67kD	24	TM
	N51260	Glutamine-fructose-6-phosphate transaminase	24	TM
RC_N51260_s	N69352	Human mRNA for KIAA0240 gene partial cds	2 4	other
RC_N69352	R37778	Homo sapiens mRNA for ATP-dependent RNA helicase #46 complete cds ESTs	24	other
RC_R37778 S82597_ma1	S82597		24	other
_	T25896	H.sapiens mRNA for UDP-GalNAc polypeptide N-acetylgalactosaminyltransferase (T1) ESTs	24	?
RC_T25896			2 4	other
RC_T29681_f	T29681	Human serine kinase mRNA complete cds	24	other
U73477	U73477	HLA-DR ASSOCIATED PROTEIN I	24	other
RC_W31919	W31919	EST CONTRACTOR OF THE PROPERTY	24	olher
W79060	W79060	ESTs Highly similar to ribosome-binding protein p34 [R norvegicus]	24	other
X13482	X13482	U2 SMALL NUCLEAR RIBONUCLEOPROTEIN A'	2 4	other
X60382_ma1	X60382	COL10A1	2 4	7
X84373	X84373	NUCLEAR FACTOR RIP140	24	other
RC_Z40898	Z40898	ESTs	2 4	other
RC_AA010065_s	AA010065	CDC28 protein kinase 2	23	other
RC_AA024658	AA024658	ESTs	23	SS,TM
RC_AA031814	AA031814	ESTs Weakly similar to R01H10 8 (C.elegans)	2 3	?
RC_AA037410_s	AA037410	Human DNA sequence from PAC 127B20 on chromosome 22q11 2-qter contains GTPase-activating protein similar to rhoGAP protein. ribosomal protein L6 pseudogene ESTs and CA repeat	23	other
RC_AA037657_s	AA037657	ESTs	23	TM
RC_AA069285	AA069285	ESTs Weakly similar to PROBABLE UBIQUITIN CARBOXYL-TERMINAL HYDROLASE R10E11.3 [C elegans]	23	other
RC_AA173223	AA173223	ESTs	2.3	other
RC_AA236951_s	AA236951	ESTs	23	other
RC_AA250737	AA250737	ESTs	2.3	other
RC_AA251776	AA251776	ESTs	23	other
RC_AA282568	AA282568	ESTs Weakly similar to F25H2 6 [C elegans]	23	other
AA330771_s	AA330771	Human protein-tyrosine phosphatase (HU-PP-1) mRNA partial sequence	23	TM
RC_AA425749	AA425749	ESTs	23	TM
RC_AA428647	AA428647	ESTs	23	other
RC_AA450116	AA450116	ESTs	23	other

FIGURE 7 (cont.)
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RC_AA459673	AA459673	ESTs Highly similar to CHROMOSOME SEGREGATION PROTEIN CUT3 [Schizosaccharomyces pombe]	23	other
RC_AA464423	AA464423	ESTs Weakly similar to !!!! ALU SUBFAMILY J WARNING ENTRY !!!! [H sapiens]	23	other
D00591	D00591	Chromosome condensation 1	2.3	?
D26156	D26156	Human mRNA for transcriptional activator hSNF2b complete cds	2.3	other
RC_F02907	F02907	ESTs	23	TM
J05633	J05633	Integrin beta-5 subunit	2.3	SS,TM
RC_N29888	N29888	Human NAD+-specific isocitrate dehydrogenase beta subunit precursor mRNA nuclear gene encoding mitochondrial protein complete cds	23	other
RC_N33063	N33063	ESTs Highly similar to GAG POLYPROTEIN [Avian spleen necrosis virus]	23	other
RC_N49284_s	N49284	MYB PROTO-ONCOGENE PROTEIN	2.3	other
RC_N66857	N66857	ESTs	2.3	?
RC_N94581	N94581	ESTs	2.3	TM
RC_R52088	R52088	E\$T - RC_R52088	2.3	other
RC_R63652	R63652	ESTs .	2.3	other
R81830	R81830	Homo sapiens breast cancer putative transcription factor (ZABC1) mRNA complete cds	2.3	other
S79873	S79873	Lysosomal-associated membrane protein 2	23	SS,TM
RC_T87807_s	T87807	ESTs	2.3	other
U37022_ma1	U37022	Human cyclin-dependent protein kinase mRNA complete cds	23	?
U47077	U47077	Human DNA-dependent protein kinase catalytic subunit (DNA-PKcs) mRNA complete cds	2.3	TM
U59423	U59423	Human chromosome 4 Mad homolog Smad1 mRNA complete cds	2.3	other
U72514	U72514	Human C2f mRNA complete cds	23	other
U77180	U77180	Human mRNA for EBI1-ligand chemokine complete cds	23	TM
RC_W23625_s	W23625	ESTS Highly similar to HYPOTHETICAL TRP-ASP REPEATS CONTAINING PROTEIN	2.3	SS,TM
		IN PMT6-PCT1 INTERGENIC REGION [Saccharomyces cerevisiae]	22	othor
RC_W49574	W49574	ESTs Moderately similar to !!!! ALU SUBFAMILY SQ WARNING ENTRY !!!! [H.sapiens]	2.3	other
X54199	X54199	Phosphoribosylglycinamide formyltransferase phosphoribosylglycinamide synthetase phosphoribosylaminoimidazole synthetase	2.3	?
X94453	X94453	Pyrroline-5-carboxylate synthetase (glutamate gamma-semialdehyde synthetase)	23	other
RC_Z39909	Z39909	ESTs Moderately similar to ZINC FINGER PROTEIN ZFP-36 [Homo sapiens]	2.3	other
Z48042	Z48042	H, sapiens mRNA encoding GPI-anchored protein p137	23	other
RC_AA011556	AA011556	ESTs	22	other
RC_AA028028	AA028028	ESTs	2 2	other
AA043160	AA043160	ESTs	22	other
RC_AA043353_s	AA043353	ESTS Highly similar to UBIQUITIN-CONJUGATING ENZYME E2-17 KD 11 [Arabidopsis	22	other
RC_AA053636	AA053636	thaliana] ESTs	22	other
RC_AA059214	AA059214	ESTs Moderately similar to neurexophilin 2 [M.musculus]	2.2	SS,
RC_AA076328_s	AA076328	Cyclin-dependent kinase inhibitor 2A (melanoma p16 inhibits CDK4)	22	TM
RC_AA126719	AA126719	ESTs	22	other
RC_AA131692	AA131692	ESTs	22	other
RC_AA148516	AA148516	ESTs .	22	other
RC_AA150088	AA150088	Homo sapiens vesicle transport related protein mRNA partial cds	22	MT
RC_AA227856	AA227856	H sapiens mRNA for HOXC9 protein exon 1	22	other
RC_AA236516	AA236516	ESTs Weakly similar to ISOLEUCYL-TRNA SYNTHETASE MITOCHONDRIAL	22	other
RC_AA251766	AA251766	[S cerevisiae] ESTs Moderately similar to metastasis-associated gene [H sapiens]	22	other
RC_AA280588	AA280588	ESTs	22	other
RC_AA287320	AA287320	ESTs	22	other
RC_AA287833	AA287833	ESTs	2.2	other
RC_AA397921	AA397921	Homo sapiens mRNA transcriptional unit N143	2.2	other
RC_AA416735	AA416735	ESTs	2.2	TM
RC_AA417030	AA417030	Homo sapiens protein regulating cytokinesis 1 (PRC1) mRNA complete cds	2.2	other
RC_AA423827_f	AA423827	ESTs	22	other
RC_AA430726	AA430726	EST - RC_AA430726	2.2	SS,
RC_AA436477	AA436477	ESTs	22	TM
RC_AA436613	AA436613	ESTs	22	other
RC_AA446949	AA446949	ESTs	22	other
RC_AA485223	AA485223	ESTs	22	TM
RC_AA490237	AA490237	EST - RC_AA490237	22	other
RC_AA495924	AA495924	ESTs	22	other
RC_AA600200	AA600200	ESTs	2.2	SS,
RC_D80237_s	D80237	Homo sapiens Arp2/3 protein complex subunit p20-Arc (ARC20) mRNA complete cds	22	?
RC_F09328	F09328	ESTs	2 2	other
RC_F13690_s	F13690	ESTs Weakly similar to ZNF127-Xp [H sapiens]	22	other
RC_H28428	H28428	ESTs	22	other
RC_H84658_s	H84658	ESTs	2 2	other

FIGURE 7 (cont.)
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RC_H99261_s	H99261	Human DNA from overlapping chromosome 19 cosmids R31396 F25451 and R31076 containing COX6B and UPKA genomic sequence	2.2	TM
RC_N39148	N39148	ESTs	22	other
RC_N90401	N90401	ESTs	22	TM
RC_N93618	N93618	ESTs	22	other
RC_N94606	N94606	ESTs	22	other
RC_R72008	R72008	ESTs Weakly similar to Diff33 gene product [H.sapiens]	22	other
R78119	R78119	ESTs	2.2	other
RC_T10060	T10060	ESTs	2.2	TM
RC_T15674_f	T15674	ESTs	22	7
RC_T59686_s	T59686	ESTs	2.2	other
U09510	U09510	Glycyl-IRNA synthetase	2.2	other
U09770	U09770	Human cysteine-rich heart protein (hCRHP) mRNA complete cds	2.2	SS,TM
U86782	U86782	Human 26S proteasome-associated pad1 homolog (POH1) mRNA complete cds	2.2	other
X70476	X70476	COATOMER BETA' SUBUNIT	2.2	?
RC_AA018587	AA018587	ESTs Weakly similar to !!!! ALU SUBFAMILY SP WARNING ENTRY !!!! [H.sapiens]	2.1	?
RC_AA134063	AA134063	ESTs	2.1	other
RC_AA158132	AA158132	ESTs Highly similar to YSA1 PROTEIN [Saccharomyces cerevisiae]	2.1	other
RC_AA251909	AA251909	Homo sapiens MAD3-like protein kinase mRNA complete cds	2.1	other
RC_AA253031	AA253031	Homo sapiens RRM RNA binding protein Gry-rbp (GRY-RBP) mRNA complete cds	2.1	other
RC_AA281780	AA281780	ESTs Weakly similar to HYPOTHETICAL 46.4 KD PROTEIN T16H12.5 IN	2.1	other
KC_74201700	201100	CHROMOSOME III [C.elegans]	2.1	omei
RC_AA291137	AA291137	ESTs	2.1	other
RC_AA393695	AA393695	LARGE FIBROBLAST PROTEOGLYCAN PRECURSOR	21	other
RC_AA400820_s	AA400820	ESTs	21	other
RC_AA403121	AA403121	ESTs	2.1	SS,
RC_AA426060	AA426060	ESTs	21	other
RC_AA427662	AA427662	ESTs	2.1	other
RC_AA451712	AA451712	ESTs	21	other
AA460077	AA460077	ESTs	2.1	other
RC_AA465148	AA465148	ESTs	2.1	other
RC_AA609869	AA609869	ESTs	21	other
RC_AA610039	AA610039	ESTs	2 1	other
RC_AA620464	AA620464	Human mRNA for KIAA0331 gene complete cds	21	2
D13988	D13988	Homo sapiens mRNA for GDP dissociation inhibitor beta	21	TM
RC_F01986_f	F01986	EST	21	2
RC_H38246_s	H38246	ESTs Weakly similar to similar to S cerevisiae LAG1 [C elegans]	21	TM
RC_H80737_s	H80737	ESTs	2 1	TM
M30938	M30938	ATP-DEPENDENT DNA HELICASE II 86 KD SUBUNIT	21	other
M74099	M74099	Cut (Drosophila)-like 1 (CCAAT displacement protein)	2.1	2
RC_N22222	N22222	ESTs	21	other
RC_N24968	N24968	Homo sapiens vacuolar H(+)-ATPase subunit mRNA complete cds	21	other
RC_N64378	N64378	ESTs	21	other
RC_N72113	N72113	ESTs	21	other
RC_N95837	N95837	Homo sapiens clone 24651 mRNA sequence	21	TM
RC_R91380_s	R91380	H sapiens RNA for CLCN3	21	TM
U51205	U51205	Human COP9 homolog (HCOP9) mRNA complete cds	21	other
U58090	U58090	Human Hs-cul-4A mRNA partial cds	21	other
U60808	U60808	Human CDP-diacylglycerol synthase (CDS) mRNA complete cds	21	TM
U61232	U61232	Human tubulin-folding cofactor E mRNA complete cds	21	other
U67122	U67122	Human ubiquitin-homology domain protein PIC1 mRNA complete cds	21	other
RC_W20391_s	W20391	Human mRNA for kinesin-related protein partial cds	21	other
RC_W32470	W32470	ESTs	2.1	other
RC_W37384_i	W37384	Homo sapiens testis-specific nm23 homolog mRNA complete cds	21	other
X70683	X70683	SRY (sex determining region Y)-box 4	21	TM
X70944	X70944	PTB-ASSOCIATED SPLICING FACTOR	21	other
RC_Z99394_s	Z99394	ESTs Moderately similar to I''' ALU SUBFAMILY SP WARNING ENTRY I''' [H sapiens]	21	other
RC_AA045481	AA045481	ESTs	2	TM
RC_AA047265	AA047265	Homo sapiens mRNA for osteoblast specific cysteine-rich protein complete cds	2	SS,
RC_AA127716	AA127716	Homo sapiens unknown mRNA complete cds	2	55, TM
RC_AA136884	AA136884	ESTs		
RC_AA181657	AA181657	ESTs	2	other
RC_AA188981	AA188981		2	other
RC_AA233177	AA233177	Homo sapiens retinoblastoma-associated protein HEC mRNA complete cds ESTs	2	? other
RC_AA237022	AA237022		2	other
110_10231022	~~231 422	ESTs	2	other

FIGURE 7 (cont.)
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RC_AA287388	AA287388	ESTs	2	other
RC_AA405838	AA405838	ESTs	2	other
RC_AA417909	AA417909	ESTs	2	other
RC_AA426375	AA426375	ESTs Highly similar to PRE-MRNA SPLICING FACTOR RNA HELICASE PRP22 [Saccharomyces cerevisiae]	2	other
AA443460	AA443460	ESTs	2	other
RC_AA443596	AA443596	ESTs	2	other
RC_AA453255	AA453255	ESTs	2	other
RC_AA476312	AA476312	ESTs	2	other
RC_AA476582	AA476582	ESTs Highly similar to SIGNAL RECOGNITION PARTICLE RECEPTOR BETA SUBUNIT [Mus musculus]	2	TM
RC_AA479139_s	AA479139	Acid phosphatase 1 soluble	2	other
RC_AA487202	AA487202	ESTs	2	other
RC_AA521474	AA521474	ESTs	2	other
RC_AA598452	AA598452	ESTs	2	?
RC_AA621122	AA621122	ESTs	2	other
AF015913	AF015913	Homo sapiens SKB1Hs mRNA complete cds .	2	other
D28364	D28364	EST - D28364	2	other
RC_D53392_f	D53392	ESTs Weakly similar to PEREGRIN [H.sapiens]	2	other
D78586	D78586	CAD PROTEIN	2	TM
D80000	D80000	Human mRNA for KIAA0178 gene partial cds	2	other
D86978	D86978	Human mRNA for KIAA0225 gene partial cds	2	TM
RC_F02651	F02651	ESTs	2	other
RC_H11938	H11938	EST - RC_H11938	2	other
RC_H78241_s	H78241	H.sapiens mRNA for novel member of serine-arginine domain protein SRrp129	2	other
L20298	L20298	Core-binding factor beta subunit	2	other
L37347	L37347	Natural resistance-associated macrophage protein 2	2	TM
M23263	M23263	Androgen receptor (dihydrotestosterone receptor testicular feminization spinal and	2	тм
RC_N22162	N22162	bulbar muscular atrophy Kennedy disease) ESTs	2	other
RC_N24954	N24954	ESTs	2	other
RC_N50963	N50963	ESTs	2	TM
RC_N70520	N70520	ESTs	2	SS,
RC_N91246	N91246	ESTs	2	2
RC_R68425	R68425	ESTs	2	,
RC_R73567	R73567	Homo sapiens meltrin-L precursor (ADAM12) mRNA complete cds	2	TM
RC_T23539	T23539	ESTs Highly similar to zinc finger protein [M musculus]	2	
T63174_s	T63174	ESTs	2	olher 2
RC_T90746	T90746	ESTs		
U05340	U05340		2	other
U34044	U34044	Human p55CDC mRNA complete cds	2	other
U37519		Human selenium donor protein (selD) mRNA complete cds	2	other
U39840	U37519	Aldehyde dehydrogenase 8	2	TM
	U39840	Human hepatocyte nuclear factor-3 alpha (HNF-3 alpha) mRNA complete cos	2	other
U91932	U91932	Human mRNA for clathrin coat assembly protein-like complete cds	2	other
W28362	W28362	ESTs	2	other
RC_W80467	W80467	ESTs	2	other
X69636	X69636	Human mRNA for KIAA0393 gene complete cds	2	other
X92896	X92896	H sapiens mRNA for ITBA2 protein	2	other
Z24724	Z24724	H.sapiens polyA site DNA	2	?
Z29090	Z29090	PHOSPHATIDYLINOSITOL 3-KINASE CATALYTIC SUBUNIT ALPHA ISOFORM	2	other
RC_Z39053	Z39053	ESTs	2	other
RC_Z40810	Z40810	ESTs	2	MT
247727	Z47727	H.sapiens mRNA for RNA polymerase II subunit	2	?
RC_AA112679	AA112679	ESTs	19	TM
AA115058_s	AA115058	ESTs	19	other
RC_AA149585	AA149585	ESTs	19	other
RC_AA173417	AA173417	ESTs	1.9	other
RC_AA227463	AA227463	ESTs Weakly similar to No definition line found [C.elegans]	1.9	2
RC_AA227963	AA227963	ESTs	1.9	¹ TM
RC_AA233168	AA233168	ESTs Highly similar to HYPOTHETICAL 16 5 KD PROTEIN IN PAS8-EGT2 INTERGENIC REGION [Saccharomyces cerevisiae]	1.9	TM
RC_AA233261	AA233261	ESTs	19	other
RC_AA236453	AA236453	ESTs	19	other
RC_AA257972	AA257972	ESTs Highly similar to UBIQUITIN-CONJUGATING ENZYME E2-17 KD 11 [Arabidopsis thaliana]	19	other
RC_AA278653_f	AA278653	ESTs	19	other
RC_AA287834	AA287834	ESTs .	19	other

FIGURE 7 (cont.)
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			1.9	7
RC_AA302745	AA302745	ESTS	1.9	?
RC_AA403008	AA403008	ESTS	1.9	7
RC_AA446918	AA446918	EST	1.9	other
RC_AA451898	AA451898	ESTs ESTs Weakly similar to Y53C12A.3 [C.elegans]	1.9	TM
AA464013	AA464013	ESTs	1.9	TM
RC_AA489046	AA489046 AA496000	ESTs	1.9	SS.
RC_AA496000	AA497052	ESTs	1.9	other
RC_AA497052	AA504832	ESTs Weakly similar to Sp140 protein [H.sapiens]	1.9	other
RC_AA504832	D12485	PLASMA-CELL MEMBRANE GLYCOPROTEIN PC-1	1.9	TM
D12485 D13645	· D13645	Human mRNA for KiAA0020 gene complete cds	1.9	other
D50920	D50920	Homo sapiens thyroid homone receptor-associated protein complex component TRAP100 mRNA complete cds	1,9	TM
H44386_s	H44386	ESTs	1.9	other
L13689	L13689	Murine leukemia viral (bmi-1) oncogene homolog	1.9	other
L33801	L33801	Human protein kinase mRNA complete cds	1.9	other
M21259	M21259	Small nuclear ribonucleoprotein polypeptide E	1.9	?
RC_N23393	N23393	ESTs	1.9	other
RC_N46423	N46423	ESTs	1.9	other
RC_N47469	N47469	ESTs	1.9	other
RC_N55336	N55336	ESTs	1.9	TM
RC_T25867	T25867	EST	1.9	?
RC_T77464	T77464	H.sapiens mRNA for transcriptional intermediary factor 2	1.9	other
RC_T89703	T89703	ESTs Weakly similar to siah binding protein 1 [H.sapiens]	1.9	other
HG174-HT174	TIGR - HG174-	EST - HG174-HT174	1.9	?
U05237	HT174 U05237	Human fetal Alz-50-reactive clone 1 (FAC1) mRNA complete cds	1.9	other
U26312	U26312	Human heterochromatin protein HP1Hs-gamma mRNA complete cds	1.9	other
U41387	U41387	Human Gu protein mRNA partial cds	1.9	other
U76638	U76638	Human BRCA1-associated RING domain protein (BARD1) mRNA complete cds	1.9	other
RC_W37933	W37933	EST - RC_W37933	1.9	?
RC_W80763	W80763	ESTs Highly similar to FK506-BINDING PROTEIN PRECURSOR [Mus musculus]	1.9	other
RC_W95063	W95063	ESTS Highly similar to HYPOTHETICAL 37.2 KD PROTEIN C12C2.09C IN CHROMOSOME I [Schizosaccharomyces pombe] Signal recognition particle 19 kD protein	1.9 1.9	TM other
X12791	X12791	H.sapiens mRNA for 2.19 gene	1.9	?
X55448_cds1	X55448	GATA-binding protein 3	1.9	other
X58072	X58072 X81788	Homo sapiens ICT1 (alias DS-1) mRNA	1.9	other
X81788	X82153	CATHEPSIN K PRECURSOR	1.9	other
X82153	Z40715	ESTs Weakly similar to T13F2.1 [C elegans]	1.9	TM
RC_Z40715 RC_AA005108	AA005108	ESTs	1.8	other
RC_AA028074	AA028074	ESTs	1.8	other
RC_AA063460_s	AA063460	Gastrin-releasing peptide	1.8	SS,
AA099241	AA099241	ESTs Moderately similar to 60S RIBOSOMAL PROTEIN L29 [H.sapiens]	1.8	other
RC_AA131584	AA131584	ESTs Weakly similar to SOF1 PROTEIN [Saccharomyces cerevisiae]	1.8	other
RC_AA191353	AA191353	ESTs	1.8	TM
RC AA232103	AA232103	ESTs	1.8	other
RC_AA232104	AA232104	ESTs Highly similar to transcription factor ARF6 chain B [M.musculus]	1.8	other
RC_AA234765	AA234765	ESTs	18	TM
RC_AA251758	AA251758	Homo sapiens spleen mitotic checkpoint BUB3 (BUB3) mRNA complete cds	1.8	other
RC_AA251982	AA251982	Homo sapiens clone 23770 mRNA sequence	1.8	other
RC_AA279171	AA279171	ESTs Weakly similar to F25D7.1 [C.elegans]	1.8	other
RC_AA283743_s	AA283743	ESTs Moderately similar to YY1-associated factor 2 [H.sapiens]	1.8	other
RC_AA291923	AA291923	ESTs	1.8	TM
RC_AA292066_i	AA292066	ESTs Weakly similar to C01H6.7 [C.elegans]	1.8	TM
RC_AA398319	AA398319	ESTs	1.8	other
RC_AA401274	AA401274	Homo sapiens RRM RNA binding protein Gry-rbp (GRY-RBP) mRNA complete cds	1.8	other
RC_AA406478	AA406478	ESTs	1.8	TM
RC_AA411144	AA411144	ESTs	1.8	TM
RC_AA417962	AA417962	ESTS Highly similar to GERANYLGERANYL PYROPHOSPHATE SYNTHETASE	1.8	other
RC_AA420988	AA420988	[Neurospora crassa] ESTs	1.8	other
RC_AA436171	AA436171	ESTs	1.8	other
RC_AA436192	AA436192	ESTs	1.8	other
RC_AA447603	AA447603	EST	1.8	?
AA455001_s	AA455001	ESTs	1.8	other

FIGURE 7 (cont.)
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00 44457500	A A 457500	FOT-	4.0	-45
RC_AA457566 RC_AA460350	AA457566 AA460350	ESTs ESTs	1.8 1.8	other other
RC_AA598988	AA598988	ESTs Moderately similar to HYPOTHETICAL 52.2 KD PROTEIN IN MPR1-GCN20	18	SS,
<del></del>		INTERGENIC REGION [Saccharomyces cerevisiae]		
RC_AA599219	AA599219 AF000430	ESTs Moderately similar to ALR [H.sapiens]  Homo sapiens mRNA for Dnm1p/Vps1p-like protein complete cds	1.8 1.8	other TM
AF000430 D13630	D13630	Human mRNA for KIAA0005 gene complete cds	1.8	other
RC_D59894	D59894	ESTs	1.8	other
RC_F02990	F02990	ESTs Highly similar to DOSAGE COMPENSATION REGULATOR [Drosophila	1.8	other
_	F04000	melanogaster]		
RC_F04989	F04989 H94248	ESTs ESTs	1.8 1.8	other other
RC_H94248 L10910	L10910	Homo sapiens splicing factor (CC1.3) mRNA complete cds	1.8	other
L25876	L25876	Human protein phosphatase (KAP1) mRNA complete cds	1.8	other
L76937_ma1	L76937	Homo sapiens Werner syndrome gene complete cds	1.8	?
M36429	M36429	Human transducin beta-2 subunit mRNA complete cds	1.8	other
M63180	M63180	Threonyl-iRNA synthetase	1.8	other
RC_N26855	N26855	ESTs Moderately similar to !!!! ALU SUBFAMILY SQ WARNING ENTRY !!!! [H.sapiens]	1.8	other
RC_N35583	N35583	ESTs Weakly similar to PROBABLE E5 PROTEIN [Human papillomavirus type 58]	1.8	?
RC_N50050	N50050	ESTs	1.8	other
RC_N52006	N52006	ESTs	1.8	TM
RC_R41281	R41281	Homo sapiens DNJ3/CPR3 mRNA complete cds	1.8	other
RC_T96595	T96595	EST - RC_T96595	1.8	TM
U14518	U14518	Centromere protein A (17kD)	1.8	other
U32986	U32986	Damage-specific DNA binding protein 1 (127 kD)	1.B	TM
U65928	U65928	V-jun avian sarcoma virus 17 oncogene homolog	1.8	other
U70322	U70322	Human transportin (TRN) mRNA complete cds	1.8	other
U72263	U72263	Exostoses (multiple) 2	1.8	TM
RC_W52225	W52225	ESTs	1.8	other
W68502	W68502 W72876	ESTs ESTs	1.8	other
RC_W72876 RC_W84790_s	W84790		1.8	SS, ?
RC_W88983	W88983	Human mRNA for KIAA0208 gene complete cds  Human RNA-binding protein CUG-BP/hNab50 (NAB50) mRNA complete cds	1.8 1.8	other
X65488	X65488	HETEROGENOUS NUCLEAR RIBONUCLEOPROTEIN U	1.8	other
X75962	X75962	OX40L RECEPTOR PRECURSOR	1.8	SS,TM
X92098	X92098	H.sapiens mRNA for transmembrane protein mp24	1.8	SS,TM
RC_Z40332	Z40332	Homo sapiens mRNA for p115 complete cds	1.8	other
RC_AA035143	AA035143	ESTs	1.7	other
RC_AA056249	AA056249	Collagen type IV alpha 3	1.7	other
RC_AA056588	AA056588	ESTs	1.7	other
RC_AA111879	AA111879	EST	1.7	7
RC_AA116075	AA116075	ESTs	1.7	other
RC_AA132514	AA132514	Homo sapiens drp1 mRNA complete cds	1.7	other
RC_AA156142_s	AA156142	ESTs	1.7	TM
RC_AA171529	AA171529	ESTs	1.7	TM
RC_AA180321	AA180321	ESTs Weakly similar to W04D2.6 [C.elegans]	1.7	other
RC_AA232315 RC_AA234767	AA232315 AA234767	Homo sapiens clone 23797 and 23917 mRNA partial cds ESTs	1.7	other
RC_AA262957	AA262957	ESTs	1.7	TM TM
RC_AA280687	AA280687	ESTs	1.7 1.7	other
RC_AA286891	AA286891	ESTs	1.7	other
RC_AA287091 s	AA287091	ESTs Highly similar to C10 [H.sapiens]	1.7	other
RC_AA291260	AA291260	ESTs	1.7	other
RC_AA400080	AA400080	EST	1.7	?
RC_AA410894	AA410894	ESTs	1.7	other
RC_AA410972	AA410972	ESTs	1.7	other
RC_AA416733	AA416733	ESTs	1.7	TM
RC_AA421773	AA421773	ESTs	1.7	other
RC_AA425439	AA425439	ESTs	1.7	other
RC_AA453465	AA453465	ESTs	1.7	other
RC_AA459005	AA459005	ESTs	1.7	other
RC_AA465690_s	AA465690	Human arginine-rich nuclear protein mRNA complete cds	1.7	other
RC_AA470140	AA470140	ESTs	1.7	?
RC_AA479362	AA479362 AA479961	ESTS	1.7	SS,
RC_AA479961	AA479961	ESTS ELCLIPE = (	1.7	other

FIGURE 7 (cont.)
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RC_AA598447	AA598447	Homo sapiens exportin t mRNA complete cds	1.7	other
RC_AA599267	AA599267	EST - RC_AA599267	1.7	other
RC_AA609053	AA609053	ESTs	1.7	SS,
RC_AA609364	AA609364	EST	1.7	7
AF002668	AF002668	Homo sapiens putative fatty acid desaturase MLD mRNA complete cds	1.7	TM
D31161_s	D31161	ESTS	1.7	TM
D78151	D78151	H.sapiens mRNA for 55.11 binding protrin	1.7	TM
RC_H15436	H15436	ESTS	1.7	other
RC_H17620	H17620	ESTs	1.7	TM
RC_H23230	H23230	ESTs	1.7	other
RC_H73608_s	H73608	ESTs	1.7	other
L19161	L19161	TRANSLATIONAL INITIATION FACTOR 2 GAMMA SUBUNIT	1.7	other
L27706	L27706	Chaperonin containing T-complex subunit 6	1.7	?
L76703	L76703	Homo sapiens protein phosphatase 2A B56-epsilon (PP2A) mRNA complete cds	1.7	?
RC_N31598	N31598	ESTS	1.7	SS,
RC_N31952	N31952	ESTs Moderately similar to HYPOTHETICAL 66.5 KD PROTEIN F02A9.5 IN CHROMOSOME III [Caenorhabditis elegans]	1.7	other
RC_N50831	N50831	ESTs	1.7	TM
RC_N51226	N51226	ESTs	1.7	other
RC_N58561_s	N58561	Cathepsin B	1.7	other
RC_N90029	N90029	Homo sapiens clone 1400 unknown protein mRNA partial cds	1.7	?
RC_N92860_s	N92860	Proto-oncogene AML1 (alternative products)	1.7	other
RC_R39923	R39923	ESTs	1.7	TM
RC_R93068	R93068	ESTs	1.7	other
RC_T03865	T03865	ESTs	1.7	other
RC_T57317	T57317	ESTs	1.7	?
HG4557-HT4962	HT4962	EST - HG4557-HT4962	1.7	?
U90551	U90551	Human histone 2A-like protein (H2AII) mRNA complete cds	1.7	other
U95367	U95367	Human GABA-A receptor pi subunit mRNA complete cds	1.7	TM
RC_W19222	W19222	ESTS	1.7	other
W23469	W23469	Homo sapiens vesicle trafficking protein sec22b mRNA complete cds	1.7	other
RC_W38150	W38150	EST - RC_W38150	1.7	?
W55890	W55890	Human Chromosome 16 BAC clone CIT987SK-A-735G6	1.7	other
RC_W85888	W85888	ESTs	1.7	other
RC_AA026418	AA026418	ESTs	1.6	other
RC_AA099589_s	AA099589	Homo sapiens mRNA for GDP dissociation inhibitor beta	1.6	TM
RC_AA101811	AA101811	EST	1.6	SS,
RC_AA121127	AA121127	ESTs Weakly similar to ZK1058.4 [C elegans]	16	SS,TM
RC_AA148885	AA148885	ESTS EST	1.6	?
RC_AA151708 RC_AA155803	AA151708 AA155803	ESTs	1.6	other
RC_AA167375	AA167375	•	1.6	other
RC_AA167708	AA167708	Homo sapiens mRNA for KIAA0530 protein partial cds ESTs	1.6	other
RC_AA181580_s	AA181580	Homo sapiens importin beta subunit mRNA complete cds	1.6 1.6	other
AA187579	AA187579	ESTs Weakly similar to Yel007c-ap [S.cerevisiae]	1.6	other
RC_AA243007	AA243007	ESTs	1.6	?
RC_AA243052	AA243052	ESTS Highly similar to GONADOTROPIN-RELEASING HORMONE RECEPTOR	1.6	other
RC_AA252360	AA252360	[Raftus norvegicus]	1.6	?
AA256106	AA256106	ESTs	1.6	other
RC_AA256678	AA256678	ESTs Highly similar to POP2 PROTEIN [Saccharomyces cerevisiae]	1.6	other
RC_AA258205	AA258205	Homo sapiens DNA polymerase zeta catalytic subunit (REV3) mRNA complete cds	1.6	other
RC_AA279667_s	AA279667	Cathepsin B	1.6	other
RC_AA347967	AA347967	ESTs	1.6	other
RC_AA417970	AA417970	ESTs	1.6	SS,
RC_AA424524	AA424524	Homo sapiens mRNA for KIAA0286 gene partial cds	1.6	?
AA426176	AA426176	ESTs Weakly similar to Similar to Sicerevisiae hypothetical protein L3111 [H.sapiens]	1.6	other
RC_AA456437	AA456437	ESTs Weakly similar to CLEAVAGE STIMULATION FACTOR 64 KD SUBUNIT	1.6	other
RC_AA456598	AA456598	[H.sapiens] ESTs	1.6	other
RC_AA463195	AA463195	ESTs	1.6	other
RC_AA465222	AA465222	ESTs	1.6	TM
RC_AA521186	AA521186	ESTs	1.6	TM
RC_AA599622	AA599622	ESTs	1.6	other
AB002343	AB002343	Human mRNA for KIAA0345 gene complete cds	1.6	TM

FIGURE 7 (cont.)
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			4.6	other
D14811	D14811	Human mRNA for KIAA0110 gene complete cds	1.6 1.6	TM
D50840	D50840	Human mRNA for ceramide glucosyltransferase complete cds	1.6	other
RC_D60856_f	D60856	Homo sapiens UDP-glucose dehydrogenase (UGDH) mRNA complete cds	1.6	other
RC_F03738_f	F03738	ESTS	1.6	TM
H19378	H19378	ESTs Homo sapiens KIAA0442 mRNA partial cds	1.6	other
RC_H95039	H95039	NAD(P)H:menadione oxidoreductase	1.6	other
J03934	J03934	Tumor protein p53 (Li-Fraumeni syndrome)	1.6	7
M22898	M22898	PROBABLE 26S PROTEASE SUBUNIT TBP-1	1.6	other
M34079	M34079 M97856	Nuclear autoantigenic sperm protein (histone-binding)	1.6	other
M97856	N26259	ESTs Weakly similar to NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 2	1.6	?
RC_N26259	N20235	[Paramecium tetraurelia]	4.6	7
RC_N69014_s	N69014	Homo sapiens SOX22 protein (SOX22) mRNA complete cds	1.6 1.6	other
RC_N73865	N73865	ESTs Weakly similar to L8004.7 gene product [S.cerevisiae]	1.6	?
RC_R10720	R10720	EST	1.6	other
RC_R15743	R15743	ESTs	1.6	?
R70621	R70621	ESTs Highly similar to hypothetical protein 100K [R.norvegicus]	1.6	other
RC_T23820	T23820	Homo sapiens cyclin T2a mRNA complete cds	1.6	?
RC_T64937_s	T64937	Homo sapiens thyroid receptor interactor (TRIP3) mRNA 3' end of cds	1.6	other
U05237	U05237	Human fetal Alz-50-reactive clone 1 (FAC1) mRNA complete cds	1.6	other
U09820	U09820	X-LINKED HELICASE II	1.6	TM
U10323	U10323	Human nuclear factor NF45 mRNA complete cds	1.6	other
U12424_s	U12424	Glycerol-3-phosphate dehydrogenase 2 (milochondrial)  Homo sapiens interferon stimulated T-cell alpha chemoattractant precursor mRNA	1.6	SS,
U59286	U59286	complete cds		
U61145	U61145	Human enhancer of zeste homolog 2 (EZH2) mRNA complete cds	1.6	other
U76992	U76992	Human Tat-SF1 mRNA complete cds	1.6	other
U90549	U90549	Human non-histone chromosomal protein (NHC) mRNA complete cds	1.6	other
U90909	U90909	Human clone 23722 mRNA sequence	1.6	other
RC_W04698	W04698	ESTs	1.6	other
RC_W15528	W15528	ESTs	1.6	other
W58247_s	W58247	ESTs Highly similar to KINESIN-LIKE PROTEIN KIF4 [Mus musculus]	1.6	other
RC_W73820	W73820	ESTs	1.6	other
X53793	X53793	MULTIFUNCTIONAL PROTEIN ADE2	1.6 1.6	other other
X58521	X58521	NUCLEAR PORE GLYCOPROTEIN P62	1.6	TM
X69910	X69910	H.sapiens p63 mRNA for transmembrane protein	1.6	other
X89059	X89059	H.sapiens mRNA for unknown protein expressed in macrophages	1.6	other
RC_Z38919	Z38919	ESTs	1.5	other
RC_AA041551	AA041551	ESTS	1.5	other
RC_AA056735	AA056735	ESTs Weakly similar to HYPOTHETICAL PROTEIN KIAA0079 [H.sapiens]	1.5	other
AA113913	AA113913	EST - AA113913	1.5	other
RC_AA133309	AA133309	EST	1.5	other
AA146888_s	AA146888	ESTs Highly similar to COATOMER ZETA SUBUNIT [Bos taurus]	1.5	other
AA195179_s	AA195179	ESTs ESTs	1.5	other
RC_AA219699 RC_AA226922	AA219699 AA226922	ESTs Highly similar to CLATHRIN COAT ASSEMBLY PROTEIN AP47 [Mus musculus]	1.5	7
RC_AA232644_s	AA232644	Protein tyrosine phosphatase non-receptor type 4	1.5	other
RC_AA236672	AA236672	ESTs Weakly similar to DFS70 [H.sapiens]	1.5	?
RC_AA256492	AA256492	ESTs	1.5	other
RC_AA262942	AA262942	ESTs	1.5	other
RC_AA279757	AA279757	ESTs Weakly similar to similar to mouse MMR1 [C.elegans]	1.5	other
RC_AA293568	AA293568	ESTs	1.5	other
RC_AA399550	AA399550	ESTs	1.5	other
RC_AA400271	AA400271	ESTs Highly similar to CALCIUM-TRANSPORTING ATPASE 1 [Saccharomyces	1.5	TM
		cerevisiae)	1.5	other
RC_AA412528	AA412528	ESTs Weakly similar to ORF2 consensus sequence encoding endonuclease and reverse transcriptase minus RNaseH [R.norvegicus]		
RC_AA433925	AA433925	ESTs	1.5	TM
RC_AA447970	AA447970	EST	1.5	TM
RC_AA476319	AA476319	ESTs .	1.5	SS,
RC_AA482014	AA482014	H.sapiens mRNA for centrin gene	1.5	other
RC_AA489086	AA489086	ESTs	1.5	other
RC_AA496257	AA496257	ESTs Weakly similar to DIPEPTIDYL PEPTIDASE IV [H.sapiens]	1.5	other
RC_AA609738	AA609738	ESTs	1.5	other
RC_AA621580	AA621580	ESTs Highly similar to HYPOTHETICAL 66.5 KD PROTEIN IN ADE12-RAP1	1.5	other
D31764	D31764	INTERGENIC REGION [Saccharomyces cerevisiae] Human mRNA for KIAA0064 gene complete cds	1.5	other

FIGURE 7 (cont.)
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D00504	B04504	No. of the ACCOUNT was partial and	4.5	albad
D38521 RC_D51177	D38521 D\$1177	Human mRNA for KIAA0077 gene partial cds ESTs	1.5 1.5	other TM
D85418	D85418	Human mRNA for phosphatidylinositol-glycan-class C (PIG-C) complete cds	1.5	TM
L18960	L18960	Eukaryotic translation initiation factor 4C (eIF-4C)	1.5	other
L33881	L33881	Protein kinase C iota	1.5	?
M31523	M31523	Transcription factor 3 (E2A immunoglobulin enhancer binding factors E12/E47)	1.5	other
M63167	M63167	V-akt murine thymoma viral oncogene homolog 1	11.5	other
RC_N21978	N21978	ESTs	1.5	other
RC_N26101	N26101	ESTs Weakly similar to DPY-30 PROTEIN (C.elegans)	1.5	other
RC_N37065	N37065	ESTs	1.5	other
RC_N48677	N48677	ESTs	1.5	TM
RC_N52271	N52271	Homo sapiens LIM protein mRNA complete cds	1.5	other
RC_N54450_i	N54450	ESTs	1.5	?
RC_N67390	N67390	ESTs	1.5	TM
RC_N68640	N68640	ESTs	1.5	other
RC_N78717_s	N78717	H.sapiens mRNA for translin	1.5	?
RC_R07016	R07016	ESTs	1.5	other
RC_R87660	R87660	EST - RC_R87660	1.5	TM
RC_T10258	T10258	EST	1.5	?
RC_T98843	T98843	ESTs Moderately similar to !!!! ALU SUBFAMILY J WARNING ENTRY !!!! [H.sapiens]	1.5	TM
HG884-HT884	TIGR - HG884- HT884	EST - HG884-HT884	1.5	7
U09564	U09564	Human serine kinase mRNA complete cds	1.5	other
U35451	U35451	Homo sapiens heterochromatin protein p25 mRNA complete cds	1.5	?
U41668	U41668	Deoxyguanosine kinase	1.5	other
U50939	U50939	Human amyloid precursor protein-binding protein 1 mRNA complete cds	1.5	other
U94836	U94836	Human ERPROT 213-21 mRNA complete cds	1.5	other
W28366	W28366	Homo sapiens clone 24800 mRNA sequence	1.5	other
RC_W72138	W72138	Homo sapiens putative transcriptional repressor E2F-6 mRNA partial cds	1.5	other
RC_W93640	W93640	ESTs	1.5	other
RC_Z39211	Z39211	Homo sapiens GDP-L-fucose pyrophosphorylase (GFPP) mRNA complete cds	1.5	other
RC_Z39255_f	Z39255	ESTs	1.5	other
RC_AA025086	AA025086	ESTs	1.4	other
RC_AA057193	AA057193	ESTs	1.4	other
RC_AA085918	AA085918	H.sapiens HUNKI mRNA	1.4	other
RC_AA114250_s	AA114250	Homo sapiens mRNA for KIAA0512 protein complete cds	1.4	other
RC_AA135095	AA135095	Homo sapiens Sox-like transcriptional factor mRNA complete cds	1.4	other
RC_AA156542	AA156542	ESTs	1.4	other
RC_AA171939	AA171939	ESTs	1.4	other
RC_AA195515	AA195515	ESTs	1.4	TM
RC_AA255554	AA255554	ESTs	1.4	TM
RC_AA262943	AA262943 AA278755	ESTs	1.4 1.4	other
RC_AA278755 RC_AA279991	AA279991	ESTs Weakly similar to !!!! ALU SUBFAMILY SB1 WARNING ENTRY !!!! [H.sapiens] ESTs	1.4	other other
AA285277	AA285277	Homo sapiens brain expressed ring finger protein mRNA complete cds	1.4	other
RC_AA287138	AA287138	ESTs Weakly similar to ASPARTYL-TRNA SYNTHETASE [Thermus aquaticus	1.4	other
-		thermophilus]		
RC_AA287879	AA287879	ESTs Highly similar to GTP-BINDING PROTEIN SARA [Mus musculus]	1.4	?
RC_AA292128	AA292128	ESTS	1.4	other
RC_AA400093	AA400093	ESTs Weakly similar to HYPOTHETICAL 48.8 KD PROTEIN IN TRK2-MRS4 INTERGENIC REGION [Saccharomyces cerevisiae]	1.4	other
AA402937	AA402937	ESTs	1.4	other
RC_AA411882	AA411882	ESTs	1.4	other
RC_AA417895	AA417895	ESTs .	1.4	SS,
AA422160	AA422160	H.sapiens NAP (nucleosome assembly protein) mRNA complete cds	1.4	other
RC_AA425100	AA425100	ESTs	1.4	other
RC_AA449068	AA449068	ESTs	1.4	TM
AA452724	AA452724	Homo sapiens TFAR19 mRNA complete cds	1.4	other
RC_AA460246	AA460246	ESTs Weakly similar to similar to tyrosyl-tRNA synthetase. [C.elegans]	1.4	other
RC_AA490949	AA490949	ESTs	1.4	other
RC_AA497015	AA497015	Homo sapiens chromosome 19 cosmid R32469	1.4	?
AB004884	AB004884	Homo sapiens mRNA for PKU-alpha partial cds	1.4	other
D38498_1	D38498	Human PMS5 mRNA (yeast mismatch repair gene PMS1 homologue) partial cds (C- terminal region)	1.4	?
RC_D80921_s	D80921	Homo sapiens clone 23965 mRNA sequence	1.4	other
RC_F04982	F04982	ESTs	1.4	other

FIGURE 7 (cont.)
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RC_F09963	F09983	ESTs	1.4	other
H59417_s	H59417	ESTs	1.4	other
RC_H93708_s	H93708	CLEAVAGE SIGNAL-1 PROTEIN	1.4	other
L06419	L06419	Lysyl hydroxylase	1,4	SS,
M29580	M29580	Zinc finger protein 7 (KOX 4 clone HF.16)	1.4	other
M62810	M62810	Transcription factor 6-like 1 (mitochondrial transcription factor 1-like)	1.4	other
RC_N23972	N23972	ESTs	1.4	other
RC_N26722	N26722	ESTs	1.4	other
RC_N64244	N64244	ESTs	1.4	TM
RC_N66569	N66569	ESTs	1.4	?
RC_N92293	N92293	EST	1.4	?
RC_R01243	R01243	ESTS	1.4	other
RC_R09196	R09196	ESTs Moderately similar to M-phase phosphoprotein 11 [H.sapiens]	1.4	other
RC_R63925	R63925	ESTS	1.4	other
RC_R64660	R64660	ESTs	1.4	?
RC_T16226	T16226	ESTS	1.4	other
RC_T17440_f	T17440	ESTs	1.4	other
U07418	U07418	DNA mismatch repair protein MLH1	1.4	other
U12595	U12595	Human tumor necrosis factor type 1 receptor associated protein (TRAP1) mRNA partial cds	1.4	other
U26727	U26727	Cyclin-dependent kinase inhibitor 2A (melanoma p16 inhibits CDK4)	1.4	TM
U84720	U84720	Homo sapiens mRNA export protein (RAE1) mRNA complete cds	1.4	other
RC_W60473	W60473	ESTs ·	1.4	other
RC_W90146_f	W90146	ESTs	1.4	other
RC_W93379_s	W93379	H.sapiens nek2 mRNA for protein kinase	1.4	other
RC_Z38501	Z38501	ESTs Weakly similar to PROBABLE E5 PROTEIN [Human papillomavirus type 58]	1.4	other
RC_Z40041	Z40041	ESTs .	1.4	other
RC_AA001386	AA001386	EST	1.3	other
RC_AA007234_s	AA007234	ESTs	1.3	other
RC_AA029264_s	AA029264	EST\$	1.3	other
RC_AA031357	AA031357	ESTs .	1.3	other
RC_AA040696_s	AA040696	ESTs	1.3	other
RC_AA046619	AA046619	ESTs	1.3	other
RC_AA059051	AA059051	ESTs	1.3	other
AA059415	AA059415	ESTs Moderately similar to !!!! ALU SUBFAMILY SB WARNING ENTRY !!!! [H sapiens]	1.3	other
AA083339	AA083339	ESTs .	1 3	other
RC_AA098864	AA098864	ESTs	1.3	other
RC_AA101601	AA101601	ESTs Highly similar to Polio virus receptor protein [H.sapiens]	1.3	other
RC_AA122394	AA122394	ESTs	1.3	other
RC_AA126426_s	AA126426	Human brain secretory protein hSec10p (HSEC10) mRNA complete cds	1.3	other
RC_AA132007_f	AA132007	Down-regulator of transcription 1 TBP-binding (negative cofactor 2)	1.3	other
AA156670_s	AA156670	Homo sapiens agrin precursor mRNA partial cds	1.3	SS,
RC_AA206800	AA206800	ESTs Weakly similar to ZINC FINGER PROTEIN 135 [H.sapiens]	1.3	TM
AA234817	AA234817	ESTs	1.3	other
RC_AA236200	AA236200	ESTs	1.3	other
RC_AA252079	AA252079	Homo sapiens mRNA for dachshund protein	1.3	other
RC_AA258189	AA258189	ESTs	1.3	other
RC_AA262889_s	AA262889	ESTs	1.3	other
RC_AA278650	AA278650	ESTs	1.3	other
AA329211_s	AA329211	Homo sapiens RRM RNA binding protein Gry-rbp (GRY-RBP) mRNA complete cds	1.3	other
RC_AA338760	AA338760	ESTs	1.3	?
RC_AA398243	AA398243	ESTs Highly similar to RSP5 PROTEIN [Saccharomyces cerevisiae]	1.3	other
RC_AA400195 RC_AA417569_i	AA400195	ESTS	1.3	other
RC_AA428992	AA417569 AA428992	ESTs ESTs	1.3	TM
RC_AA435536	AA435536	ESTs	1.3	other
RC_AA443294	AA443294		1.3	other
RC_AA449071	AA449071	Homo sapiens putative transcriptional repressor E2F-6 mRNA partial cds ESTs	1.3	other
AA458542	AA4490/1 AA458542		1.3	'TM
RC_AA461169	AA461169	Homo sapiens chromosome 19 cosmid R32469 ESTs	1.3	other
RC_AA464428	AA464428	ESTs	13	other
RC_AA465093	AA465093	ESTs .	1.3	other
RC_AA485424	AA485424	ESTs	1.3	other
RC_AA487492_s	AA487492		1.3	other
	. W 1701 70Z	Homo sapiens clone 23592 mRNA sequence	1.3	other

FIGURE 7 (cont.)
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RC_AA504499	AA504499	ESTs Highly similar to probable chloride channel 3 [H.sapiens]	1.3	other
RC AA521471	AA521471	ESTs	1.3	other
RC_AA598506_s	AA598506	Human mRNA for KIAA0179 gene partial cds	1.3	other
RC_AA598675	AA598675	ESTs	1.3	other
RC_AA599718	AA599718	H.sapiens mRNA for translin associated protein X	1.3	other
RC_D11718	D11718	ESTS	1.3	?
D87466	D87466	Human mRNA for KIAA0276 gene partial cds	1.3	other
RC_F13663	F13663	ESTs	1.3	other
RC_H26417	H26417	ESTs	1.3	other
RC_H38086	H38086	Human N-ethylmaleimide-sensitive factor mRNA partial cds	1.3	other
RC_H38828_s	H38828	H.sapiens RBQ-1 mRNA	1.3	other
RC_H60061	H60061	ESTs Moderately similar to !!!! ALU SUBFAMILY SB WARNING ENTRY !!!! [H.sapiens]	1.3	other
RC_H71863_s	H71863	Zinc finger protein 139 (clone pHZ-37)	1.3	other
RC_H83438_s	H83438	Homo sapiens mRNA for DDS1beta protein complete cds	1.3	other
M64929	M64929	Protein phosphatase 2 (formerly 2A) regulatory subunit B (PR 52) alpha isoform	1.3	other
M95767	M95767	DI-N-ACETYLCHITOBIASE PRECURSOR	1.3	SS,
RC_N20630_i	N20630	ESTs	1.3	other
RC_N24732	N24732	ESTs	1.3	other
RC_N51855	N51855	ESTs Moderately similar to NAD(+) ADP-RIBOSYLTRANSFERASE [D melanogaster]	1.3	other
RC_R49886	R49886	ESTs	1.3	SS.
RC_T23932_f	T23932	ESTs	1.3	other
RC_T40707	T40707	ESTs	1.3	other
RC_T59859	T59859	ESTs	1.3	other
RC_T64438	T64438	ESTs Weakly similar to C01A2.4 [C.elegans]	1.3	TM
T68510	T68510	ESTs	1.3	other
RC_T95591	T95591	ESTs	1.3	other
U02680	U02680	Human protein tyrosine kinase mRNA complete cds	1.3	other
U28686	U28686	Human putative RNA binding protein RNPL mRNA complete cds	1.3	other
U66561	U66561	Human kruppel-related zinc finger protein (ZNF184) mRNA partial cds	1.3	other
U96113	U96113	EST - U96113	1.3	other
RC_W52065_f	W52065	Homo sapiens mRNA for KIAA0539 protein complete cds	1.3	?
RC_W67524	W67524	Human protein-tyrosine phosphatase (HU-PP-1) mRNA partial sequence	1.3	TM
RC_W86978	W86978	ESTs	1.3	other
X69398	X69398	CD47 antigen (Rh-related antigen integrin-associated signal transducer)	1.3	SS,TM
X97544	X97544	H.sapiens mRNA for TIM17 preprotein translocase	1.3	TM
RC_Z41963_r	Z41963	Homo sapiens HP protein (HP) mRNA complete cds	1.3	?
Z46629	Z46629	SRY (sex-determining region Y)-box 9 (campomelic dysplasia autosomal sex-reversal)	1.3	other
RC_AA010188	AA010188	ESTs	1.2	other
RC_AA025746	AA025746	ESTs	1.2	other
AA112222	AA112222	EST - AA112222	1.2	other
AA147543	AA147543	ESTs	1.2	SS,
AA355201	AA355201	ESTs	1.2	SS,TM
RC_AA398222	AA398222	ESTs	1.2	other
RC_AA411708	AA411708	Homo sapiens clone 23685 mRNA sequence	1.2	other
RC_AA433943	AA433943	ESTs Highly similar to 50S RIBOSOMAL PROTEIN L13 [Mycobacterium leprae]	1.2	other
RC_AA464758	AA464758	ESTs	1.2	other
RC_H05635	H05635	ESTs	1.2	TM
L38961	L38961	Integral transmembrane protein 1	1.2	MT
N42440	N42440	ESTs Weakly similar to hnRNA-binding protein M4 [H.sapiens]	1.2	other
RC_N55304_s	N55304	ESTs	1.2	other
RC_N67104	N67104	ESTs	1.2	other
RC_N68622 RC_N71027	N68622 N71027	ESTs Highly similar to HYPOTHETICAL 27.5 KD PROTEIN IN SPX19-GCR2 INTERGENIC REGION [Saccharomyces cerevisiae]	1.2	other
RC_N74635	N74635	ESTs	1.2	other
RC_R62444	R62444	ESTs	1.2	other
RC_T17498	T17498	ESTs	1.2	other
RC_T32794_s	T32794 .	ESTs	1.2	TM
RC_T85190	T85190	EST - RC_T85190	1.2	other
RC_T99364	T99364	ESTs Weakly similar to !!!! ALU SUBFAMILY J WARNING ENTRY !!!! [H.sapiens]	1.2	?
U20240	U20240	CCAAT/enhancer binding protein (C/EBP) gamma	1.2	other
U51698	U51698	ESTs	1.2 1.2	other
U79718	U79718	Human endonuclease III homolog mRNA complete cds		?
W03007	W03007	ESTs	1.2	other
			1.2	other

FIGURE 7 (cont.)
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RC_W61011	W61011	ESTs	1.2	other
RC_W87544	W87544	ESTs	1.2	other
X02751	X02751	Neuroblastoma RAS viral (v-ras) oncogene homolog	1.2	?
Z14077_s	Z14077	YY1 transcription factor	1.2	other
RC 238839	Z38839	ESTs	12	?

FIGURE 7 (cont.)
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<b>T</b> 0-	Accession	PROBESET	1117 (23.1)	11 (1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
Title	T78889	RC_T78889_s	3.1	44.4
fibronectin 1	AA393803	RC_AA393803	3.9	24.6
ESTs		RC_H88540	2.8	22.0
Human heart mRNA for heat shock protein 9; partial cds	H88540 W31478	RC_W31478_s	1.5	20.8
ESTs	VV31476	NO_4451476_5	1.5	20.0
ESTs; Weakly similar to (defline not available 4454131) [D.melanogaster]	W79424	RC_W79424_s	1.8	19.7
Homo sapiens RGS-GAIP interacting protein GIPC mRNA; complete cds	AA149940	RC_AA149940	1.2	17.5
collagen; type I; alpha 2	J03464	J03464_s	8.7	17.3
Homo sapiens mRNA for actin-related protein; complete cds	W48638	RC_W48638	3.0	17.3
MYOSIN REGULATORY LIGHT CHAIN 2; NONSARCOMERIC	W92462	RC_W92462	1.1	16.1
ESTs; Weakly similar to NADH-UBIQUINONE		_		
OXIDOREDUCTASE CHAIN 5 [Ascaris suum]	D63079	D63079_s	1.9	15.9
ESTs	AA255874	RC_AA255874	1.7	15.8
fibronectin 1	X02761	X02761	3.6	15.2
TFAR19 novel apoptosis-related gene	R71082	RC_R71082_s	2.5	15.2
ESTs; Highly similar to FRIZZLED PROTEIN PRECURSOR				
[Drosophila melanogaster]	AA449749	RC_AA449749	10.6	14.7
ESTs	AA243721	RC_AA243721	1.5	14.0
Homo sapiens mRNA for KIAA886 protein; complete cds	AA600169	RC_AA600169	1.2	13.9
ESTs	AA182001	RC_AA182001_i	1.2	13.8
dihydropyrimidinase-like 2	T10322	RC_T10322	8.0	13.5
	M97935	AFFX-HUMISGF3A/M97935_3	2.3	13.5
ESTs	AA186897	RC_AA186897	4.5	13.5
lactate dehydrogenase B	T16206	RC_T16206_i	0.6	13.1
ESTs; Weakly similar to SAS [H.sapiens]	AA365742	AA365742_s	3.2	13.0
ESTs	AA621274	RC_AA621274_i	1.1	12.8
ESTs; Highly similar to GALECTIN-1 [Homo sapiens]	H25999	RC_H25999_s	2.5	12.7
H.sapiens gene from PAC 295C6; similar to rat PO44	AA281132	RC_AA281132	1.9	12.6
Human CCAAT-box-binding factor (CBF) mRNA; complete cds	T47491	RC_T47491	1.4	12.3
ESTs; Moderately similar to 25E8.I [D.melanogaster]	W79421	RC_W79421	1.2	12.0
ESTs; Highly similar to UBIQUITIN-CONJUGATING ENZYME		DO 1170205 -	1.7	11.7
E2-25 KD [Bos taurus]	H78385	RC_H78385_s	14.8	11.5
ESTs	F13673	RC_F13673 AA247685	4.3	11.5
desmoplakin (DPI; DPII)	AA247685		1.6	11.4
immunoglobulin gamma 3 (Gm marker)	H64493	RC_H64493_f	1.4	11.4
Pantophysin [human; keratinocyte line HaCaT; mRNA; 216 nt]	R72029	RC_R72029_f	2.7	11.4
ESTs	D79891	D79891	2.0	11.3
ESTs	AA167393	RC_AA167393_s	1.9	11.2
Human transcriptional coactivator PC4 mRNA; complete cds	D57317	RC_D57317	1.3	
ESTs	N67507	RC_N67507	1.3	
neurotrophic tyrosine kinase; receptor-related 1 ESTs; Highly similar to HYPOTHETICAL 1.4 KD PROTEIN IN UBP5-SPT15 INTERGENIC REGION [Saccharomyces	C15347	RC_C15347		
cerevisiae]	AA027086	RC_AA027086	2.3	11.2
ESTs; Weakly similar to EBNA-2 NUCLEAR PROTEIN [Human herpesvirus 4 (strain b95-8)]	AA598781	RC_AA598781	4.0	11.2
Homo sapiens actin-related protein Arp2 (ARP2) mRNA; complete cds	C16379	C16379	1.5	11.1

#### FIGURE 8

tyrosine 3-monooxygenase/tryptophan 5-monooxygenase				
activation protein; eta polypeptide	H69844	RC_H69844_s	1.5	11.1
Homo sapiens mRNA for DCRA; complete cds	R97540	RC_R97540_f	1.0	11.1
Homo sapiens chaperonin containing t-complex polypeptide 1; beta subunit (Cctb) mRNA; complete cds	AA488991	RC_AA488991_s	1.5	10.9
Homo sapiens actin-related protein Arp2 (ARP2) mRNA;				
complete cds	AF006082	AF006082	1.6	10.9
ESTs	T23457	RC_T23457	3.7	10.8
Human (clone E5.1) RNA-binding protein mRNA; complete cds	T33593	RC_T33593_s	2.0	10.8
Homo sapiens clone 24416 mRNA sequence	AA417761	RC_AA417761	1.4	10.7
ESTs	N39152	RC_N39152	1.5	10.7
ESTs	AA429539	AA429539_f	1.8	10.7
ESTs	T30617	T30617	1.1	10.6
small inducible cytokine A5 (RANTES)	AA486072	RC_AA486072_i	1.4	10.6
ESTs	R79392	RC_R79392	3.3	10.5
calumenin	AA477316	RC_AA477316	2.8	10.5
ESTs	R54421	RC_R54421_s	1.4	10.4
UDP-N-acteylglucosamine pyrophosphorylase 1; Sperm				
associated antigen 2	AA447549	RC_AA447549	1.9	10.3
Homo sapiens HRIHFB2115 mRNA; partial cds	H11320	RC_H11320_s	2.0	10.3
ESTs; Moderately similar to putative G-binding protein	110.4077	DO 1104077	• • •	
[H.sapiens]	H94877	RC_H94877	1.8	10.2
protein kinase; cAMP-dependent; catalytic; alpha	H89514	RC_H89514_s	1.0	10.2
ESTs	F10354	RC_F10354_f	1.0	10.1
ESTs zn13e4.r1 Stratagene hNT neuron (#937233) Homo sapiens cDNA clone IMAGE:547326 5' similar to gb:J2683 ADP,ATP	AA173981	RC_AA173981	1.2	10.0
CARRIER PROTEIN, FIBROBLAST ISOFORM (HUMAN);, mRNA sequence	AA084874	AA084874_f	0.9	10.0
ESTs; Highly similar to YME1 PROTEIN [Saccharomyces cerevisiae]	AA452161	RC_AA452161	1.8	9.9
ESTs	N93521	RC_N93521	1.5	9.9
small inducible cytokine A5 (RANTES)	M21121	M21121_s	0.9	9.9
ESTs	AA490112	RC_AA490112_s	2.1	9.9
ESTs; Highly similar to HYPOTHETICAL 16.3 KD PROTEIN IN DUR1;2-NGR1 INTERGENIC REGION [Saccharomyces			2.1	3.3
cerevisiae]	AA053139	RC_AA053139	3.2	9.7
ESTs	AA446461	RC_AA446461	1.1	9.7
C-terminal binding protein 2	N50048	RC_N50048	2.1	9.7
Homo sapiens mRNA for putative progesterone binding protein	N66130	RC_N66130	1.4	9.6
ESTs transcription factor AP-2 alpha (activating enhancer-binding	AA490341	RC_AA490341_s	1.4	9.5
protein 2 alpha) ESTs; Highly similar to (defline not available 468665)	R38044	RC_R38044_f	9.4	9.4
[H.sapiens]	W73805	W73805	1.2	9.4
cathepsin B ESTs; Highly similar to heat shock factor binding protein 1	AA608751	RC_AA608751_i	2.1	9.3
HSBP1 [H.sapiens]	D59525	RC_D59525_f	1.8	9.3
ESTs	AA280409	RC_AA280409_s	2.1	9.3
ESTs; Weakly similar to similar to yeast adenylate cyclase [H.sapiens]	N77542	N77542	1.6	9.2
ESTs	D60296	RC_D60296	1.7	9.2
ESTs; Weakly similar to VACUOLAR ATP SYNTHASE 54 KD SUBUNIT [Saccharomyces cerevisiae]	Z39349	RC_Z39349	1.6	9.2
solute carrier family 12 (sodium/potassium/chloride				
transporters); member 2	AA262080	RC_AA262080	1.4	9.2
iduronate 2-sulfatase (Hunter syndrome)	H14810	RC_H14810_s	1.0	9.1

		,		
ESTs	T90531	RC_T90531	1.5	9.1
ESTs	Z40959	RC_Z40959_f	1.0	8.9
collagen-binding protein 2 (colligen 2)	H27188	RC_H27188_f	2.6	8.9
HMT1 (hnRNP methyltransferase; S. cerevisiae)-like 1	T81393	RC_T81393_s	1.8	8.8
Homo sapiens lysophospholipase (LPL1) mRNA; complete cds	AA252436	AA252436	1.6	8.8
Homo sapiens TACC1 (TACC1) mRNA; complete cds	N46837	RC_N46837	2.4	8.7
ESTs	AA122386	RC_AA122386	6.6	8.7
ESTs; Highly similar to LEUCYL-TRNA SYNTHETASE; CYTOPLASMIC [Saccharomyces cerevisiae]	R32993	R32993_s	1.6	8.6
NADH dehydrogenase (ubiquinone) 1 beta subcomplex; 6	046330	040000		
(17kD; B17)	C16329	C16329	0.9	8.6
ESTs; Weakly similar to transporter protein [H.sapiens] ESTs	R80048	R80048	1.2	8.5
	AA102644	RC_AA102644	1.8	8.5
ESTs; Weakly similar to (defline not available 4234) [D.melanogaster]	AA393805	RC_AA393805	1.1	8.4
ESTs; Weakly similar to putative Rab5-interacting protein {clone L1-94} [H.sapiens]	H68794	RC H68794	4 6	ا ، ،
ESTs	AA399445	RC_AA399445	1.5	8.4
	AA388445	RC_AA399445	0.9	8.4
Homo sapiens Arp2/3 protein complex subunit p16-Arc (ARC16) mRNA; complete cds	AF006088	AF006088	1.5	8.3
ESTs	AA278329	RC_AA278329_f	3.1	8.3
ESTs	AA187490	RC_AA187490	3.6	8.3
ESTs	N90933	RC_N90933	1.0	8.2
ESTs; Weakly simitar to predicted using Genefinder [C.elegans]	D31058	D31058 s	2.1	8.2
immumoglobulin lambda gene cluster	T67053	RC_T67053_f	1.2	8.2
epithelial membrane protein 2	T88721	RC_T88721_s	1.3	8.2
Homo sapiens actin-related protein Arp3 (ARP3) mRNA;				
complete cds	AF006083	AF006083	1.8	8.1
ESTs	AA040923	RC_AA040923	1.8	8.1
INTERFERON-ALPHA INDUCED 11.5 KD PROTEIN	AA161292	RC_AA161292_s	1.5	8.0
ESTs	W85875	RC_W85875	0.9	8.0
Human mRNA for KIAA336 gene; complete cds	AA608903	RC_AA608903	1.4	7.9
ESTs; Moderately similar to KIAA438 [H.sapiens]	H81379	RC_H81379_s	1.3	7.9
H. sapiens cDNA for RFG	AA194075	RC_AA194075_f	0.4	7.9
ESTs; Weakly similar to cDNA EST EMBL:T1157 comes from this gene [C.elegans]	N67312	RC_N67312	1.7	7.9
	M10098	AFFX-HUMRGE/M10098_5		
Fibronectin, Alt. Splice 1	HG3044-HT3742	HG3044-HT3742	1.1 3.0	7.9 7.8
cytochrome c oxidase subunit VII-related protein	AA025213	RC_AA025213	1.3	7.8
CD74 antigen (invariant polypeptide of major histocompatibility complex; class II antigen-associated)	W67577	RC_W67577_s	1.2	[
ESTs; Weakly similar to neural differentiation-associated protein	***************************************	110_4407077_3	1.2	7.8
[M.musculus]	AA233342	RC_AA233342	3.8	7.7
ESTs	AA291159	RC_AA291159_f	0.7	7.7
ESTs	N63604	RC_N63604	3.6	7.7
HEAT SHOCK 7 KD PROTEIN 1	T66307	RC_T66307_f	1.3	7.6
Human DNA sequence from clone 3M3 on chromosome 6p22.1-22.3. Contains three novel genes; one similar to C. elegans Y63D3A.4 and one similar to (predicted) plant; worm; yeast and archaea bacterial genes; and the first exon of the KIAA319 gene. Contains E	AA243497	RC AA243497	1.7	7.6
Homo sapiens cDNA for dihydroxyacetone phosphate		_		}
acyltransferase (DAP-AT) ESTs	AA600134	RC_AA600134	1.7	7.6
	H61476	RC_H61476_s	1.6	7.6
transforming growth factor; beta receptor II (7-8kD)	H90886	RC_H90886_s	0.8	7.6

STs; Weakly similar to ubiquitous TPR motif; Y isoform I.sapiens]	AA449320	RC_AA449320	1.0	7.5
STs; Highly similar to HYPOTHETICAL 37.7 KD PROTEIN K686.3 IN CHROMOSOME III [Caenorhabditis elegans]	N48787	RC_N48787	1.9	7.9
omo sapiens heterogeneous nuclear ribonucleoprotein R iRNA; complete cds	C16574	C16574	1.7	7.5
omo sapiens mRNA for CMP-sialic acid transporter; complete		<b>DO</b> 444-11-11		_
ds	AA481542	RC_AA481542_s	1.2	7.9
STs; Weakly similar to F15D4.3 [C.elegans]	N89563	N89563_s	2.0	7.
STs	AA490262	RC_AA490262	2.9	7.
STs; Weakly similar to similar to Yeast hypothetical protein 8167.12 like [C.elegans]	AA621349	RC_AA621349	1.3	7.
STs; Highly similar to (defline not available 412715)  1.sapiens]	C01782	C01782	1.9	7.
STs	AA402492	RC_AA402492	1.7	7.
	AFFX-M27830	AFFX-M27830_5	0.5	
STe: Washly similar to C17411 & [C alegans]		RC_AA194237	1.7	7.4 7.4
STs; Weakly similar to C17H11.6 [C.elegans] STs	AA194237 H10933	RC_AA194237 RC_H10933		
uman spliceosomal protein (SAP 49) gene; complete cds	AA463934	RC_AA463934	4.6 1.6	7.4 7.3
uman spliceosomal protein (SAP 49) gene; complete cus uman mRNA for KIAA174 gene; complete cds	R16097	RC_AA463934 RC_R16097_s	1.5	7. 7.
uman mkiva for Klaat74 gene; complete cos STs	AA128486	RC_R16097_s RC_AA128486	1.2 1.5	7. 7.
STs; Weakly similar to HYPOTHETICAL 128.5 KD HELICASE	AA120400	RO_AA120400	1.5	7.
ATS1-TPD3 INTERGENIC REGION [Saccharomyces erevisiae]	R59694	RC_R59694_s	1.4	7.
ST's	AA428090	AA428090	7.0	7. 7.
STs	H88486	RC_H88486_f	1.5	7.
STs; Highly similar to PROTEIN TRANSPORT PROTEIN	1100-100	110_1100+00_1	1.5	,
EC61 GAMMA SUBUNIT [Canis familiaris; Mus musculus]	D79052	D79052_s	3.1	7.
-terminal binding protein 2	AA417287	RC_AA417287	2.6	7.
rotein kinase; mitogen-activated 6 (extracellular gnal-regulated kinase; p97)	T32837	RC_T32837_\$	1.1	6.
mor rejection antigen (gp96) 1	D51235	RC_D51235_f	1.2	6.
STs; Highly similar to INORGANIC PYROPHOSPHATASE				
Bos taurus]	F04258	RC_F04258_s	3.3	6.
uman mRNA for KIAA35 gene; partial cds	D51272	RC_D51272_s	3.2	6.
omo sapiens mRNA from chromosome 5q31-33 region	T99196	RC_T99196_s	1.4	6.
sapiens mRNA for serine palmitoyltransferase; subunit I	T39740	T39740_s	1.3	6.9
omo sapiens calcium binding protein (ALG-2) mRNA; complete	AA122332	DC 4442222	4.7	
STs	T92245	RC_AA122332 RC_T92245_i	1.7 0.9	6.9 6.9
STs	F01813	RC_F01813_s	3.1	6.9
STs; Highly similar to putative Rab5-interacting protein {clone 1-57} [H.sapiens]	AA292533	RC_AA292533	1.3	6.9
STs; Moderately similar to POSSIBLE DNA-REPAIR ROTEIN XP-E [Cercopithecus aethiops]	AA287961	RC_AA287961		
STs	AA053883	RC_AA257951 RC_AA053883	1.6	6.8
eptidylprolyl isomerase B (cyclophilin B)	H96665	RC_H96665_s	0.7 2.2	6.8
onnective tissue growth factor	AA449789	RC_AA449789_f	1.9	6.8
omo sapiens exportin t mRNA; complete cds				6.8
STs; Weakly similar to Ydr372cp [S.cerevisiae]	H99877	RC_H99877	4.0	6.8
STs; Highly similar to (defline not available 4426962)	AA191014	RC_AA191014	1.7	6.8
ł.sapiens] STs	H82061	RC_H82061	1.2	6.8
ais	AA433947	RC_AA433947	1.8	6.8
STs	AA236280	RC_AA236280	1.5	6.8
	AA236280 AA521303	RC_AA236280 RC_AA521303	1.5 0.8	6.8 6.7

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proteasome (prosome; macropain) 26S subunit; non-ATPase; 1	AA460532	RC_AA460532	1.5	6.6
ESTs	AA398197	RC_AA398197	1.9	6.6
cytochrome c oxidase subunit IV	AA236361	RC_AA236361	1.2	6.6
Thymosin; beta 1	T59161	RC_T59161_s	2.6	6.6
Homo sapiens mRNA for KIAA733 protein; partial cds	H88033	H88033_s	1.2	6.6
ESTs	H86543	RC_H86543_f	1.8	6.6
glucan (1;4-alpha-); branching enzyme 1 (glycogen branching enzyme; Andersen disease; glycogen storage disease type IV)	H71861	RC_H71861_s	0.9	6.6
ESTs	T95333	RC_T95333	5.3	6.6
ESTs; Moderately similar to !!!! ALU CLASS C WARNING ENTRY !!!! [H.sapiens]	AA425447	RC_AA425447	1.7	6.6
Homo sapiens GA17 protein mRNA; complete cds	AA147725	RC_AA147725	2.5	6.5
lactate dehydrogenase A	AA112012	RC_AA112012_s	2.0	6.5
ESTs	AA621159	RC_AA621159	1.8	6.5
ESTs; Highly similar to PRE-MRNA SPLICING FACTOR RNA HELICASE PRP22 [Saccharomyces cerevisiae]	AA429228	AA429228	1.6	6.5
collagen; type I; alpha 2	Z74616	Z74616	9.9	6.5
ESTs; Weakly similar to !!!! ALU SUBFAMILY SX WARNING				
ENTRY !!!! [H.sapiens]	AA608668	RC_AA608668	1.0	6.5
ESTs	N93155	RC_N93155_i	1.8	6.5
ESTs	D51401	RC_D51401_s	1.4	6.3
Homo sapiens mRNA for KIAA96 protein; partial cds	AA250870	AA250870_s	2.3	6.3
ESTs	AA429636	RC_AA429636	0.9	6.3
ESTs; Weakly similar to similar to Yeast hypothetical protein L8167.12 like [C.elegans]	H73484	RC_H73484_s	1.3	6.3
ESTs	AA489091	RC_AA489091	1.4	6.3
ESTs; Highly similar to HYPOTHETICAL 29.4 KD PROTEIN IN STE6-LOS1 INTERGENIC REGION [Saccharomyces		DO 1104004 :	1.1	6.3
cerevisiae]	H84891	RC_H84891_i	1.1	6.3
ESTs	AA001049	RC_AA001049	1.1	6.3
ESTs; Weakly similar to !!!! ALU SUBFAMILY SP WARNING ENTRY !!!! [H.sapiens]	N20066	RC_N20066	1.2	6.2
ESTs; Highly similar to RAS-RELATED PROTEIN RAB-1	4.4.400070	DC 44439970	1.8	6.2
[Canis familiaris]	AA428870	RC_AA428870	2.2	6.2
ESTs	W16836	RC_W16836_s		
ESTs	H07873	RC_H07873	1.2	6.2
ESTs	W58619	RC_W58619	1.7	6.2
Homo sapiens mRNA for KIAA737 protein; complete cds	N66219	RC_N66219	1.3	6.2
bone morphogenetic protein 6	AA092596	AA092596	1.1	6.2
Homo sapiens DNA from chromosome 19-cosmid R3879 containing USF2; genomic sequence	R36881	RC_R36881_s	1.5	6.2
ESTs	AA179387	RC_AA179387	4.0	6.1
ESTs; Moderately similar to fibronectin [H.sapiens]	AA279397	RC_AA279397	1.3	6.1
ESTs; Highly similar to MYO-INOSITOL-1(OR 4)-MONOPHOSPHATASE [Xenopus laevis]	T96374	RC_T96374	0.8	6.1
ESTs	AA465194	RC_AA465194	1.7	6.1
Human amino acid transport-related protein mRNA; complete		_		Ì
cds	AA152418	RC_AA152418	1.1	6.1
ESTs	AA447971	RC_AA447971	5.1	6.1
ESTs	W38419	RC_W38419_f	0.9	6.1
pigment epithelium-derived factor	AA111889	RC_AA111889	1.5	6.1
ESTs	W42508	RC_W42508	1.1	6.1
ESTs	N91023	RC_N91023	3.3	6.1

Homo sapiens 3-phosphoglycerate dehydrogenase mRNA; compilet cols   Tas3846   RC_T83846   0.9   6.0	collagen; type I; alpha 2	H88674	RC H88674 s	3.0	6.0
Complete cds				0.0	
LYMPHOCYTE-SPECIFIC PROTEIN LSPI Homo saplens secreted cament gland protein XAG-2 homolog (NAG-ZR) miNA(: complete code Homo saplens secreted cament gland protein XAG-2 homolog (NAG-ZR) miNA(: complete code FSTs; Moderately similar to Similar to S. cerevisiae hypothetical protein L311 [H.saplens] ESTs AA06163 AA06163 RC_AA06163 1.1 5.9 ESTs ESTs AA06163 RC_AA06163 1.1 5.9 ESTs AA06163 RC_AA06163 1.1 5.9 ESTs ESTs AA06163 RC_AA06163 1.1 5.9 ESTs AA06163 RC_AA06163 1.1 5.9 ESTs ESTs AA06163 RC_AA06163 1.1 5.9 ESTs AA06163 RC_AA06163 1.1 5.9 ESTs ESTs AA06163 RC_AA06163 1.1 5.9 ESTs AA06163 RC_AA06163 1.1 5.9 ESTs ESTs AA06163 RC_AA06163 RC_AA06163 RC_AA06163 1.1 5.9 ESTs ESTs AA06163 RC_AA06163 RC_AA06173 RC_AA06163 RC_AA06163 RC_AA06163 RC_AA06163 RC_AA06173 RC_A		T83646	RC_T83646	0.9	6.0
Homo sapiens secreted carrient gland protein XAG-2 homolog (hAG-2/IR mRAV) complete cds   1.3   5.9	HMT1 (hnRNP methyltransferase; S. cerevisiae)-like 2	W46810	RC_W46810_s	3.2	5.9
AA421662   RC_AA421662   1.3   5.9	LYMPHOCYTE-SPECIFIC PROTEIN LSP1	T49291	RC_T49291_s	1.0	5.9
ESTs; Moderately similar to Similar to S. cerevisiae hypothetical protein [1411 [H.sapiens]		AA421562	RC_AA421562	1.3	5.9
Drotein L3111 [H.sapiens]	Homo sapiens clone 23956 mRNA; partial cds	W69452	RC_W69452	1.1	5.9
ESTs		N79531	RC N79531 s	3.1	5.9
ESTs	l'	AA406163	<del>-</del> -		5.9
Homo sapiens clone 2394 mRNA sequence		AA454157	<del>-</del>		i i
ESTs	ł	AA609773	<del></del>	5.5	5.9
Complete ods	ESTs	AA156897	AA156897_s	3.7	5.8
Diliverdin reductase A		AA021623	RC AA021623 s	1.1	5.8
ESTs; Weakly similar to hypothetical protein [H. sapiens] T97257 RC_T97257_f 1.4 5.8 quinoid dihydropteridine reductase T69009 RC_T69009_s 0.5 5.8 ESTs AA489459 RC_AA489459 R.1.1 5.7 ESTs; Highly similar to follistatin-related protein [H.sapiens] D51110 RC_D51110_s 2.1 5.7 ESTs; Highly similar to !!!! ALU SUBFAMILY SP WARNING ENTRY !!!! [H. sapiens] AA452855 RC_AA452855 RC_AA452855 RC_BTY, Moderately similar to !!!! ALU SUBFAMILY SP WARNING ENTRY !!!! [H. sapiens] AA442125 RC_AA452855 RC_BSTs; Highly similar to EUXARYOTIC INITIATION FACTOR 4 GAMMA [Oryctolagus cuniculus] W84870 RC_W84870_s 1.2 5.7 ESTs; Moderately similar to EUXARYOTIC INITIATION FACTOR 4 GAMMA [Oryctolagus cuniculus] AA45027 RC_AA45027 1.9 5.7 ESTs; Wooderately similar to HN1 [M.musculus] AA45027 RC_AA45027 1.9 5.7 ESTs; Wooderately similar to HN1 [M.musculus] AA45027 RC_AA45027 1.9 5.7 ESTs; Weakly similar to brain-specific L-proline transporter AA460049 AA460049_s 1.2 5.7 ESTs; Weakly similar to brain-specific L-proline transporter AA480049 AA460049_s 1.2 5.7 ESTs AA453783 RC_AA453783 RC_AA453783_s 3.7 5.6 ESTs AA453783 RC_AA453783_s RC_AA453783_s 3.7 5.6 ESTs AA53783 RC_AA453783_s RC_AA453783_s 3.7 5.6 ESTs; Weakly similar to (defline not available 446577) [H. sapiens] AA45027 RC_AA3504914 0.8 5.6 ESTs; Weakly similar to (defline not available 446577) [H. sapiens] AA469614 RC_AA48914 0.8 5.6 Globalstoma amplified sequence AA095021 AA095021 1.0 5.6 Karyopherin alpha 1 (importin alpha 5) N35247 RC_AA35461 0.8 5.6 Globalstoma amplified sequence AA095021 AA095021 1.0 5.6 Karyopherin alpha 1 (importin alpha 5) N35247 RC_A33978 RC_A348361 1.6 5.5 ESTs H98153 RC_A4453461 RC_A4453461 1.6 5.5 ESTs H98153 RC_A4453461 RC_A4453461 1.6 5.5 ESTs H98153 RC_A4453461 RC_A453461 1.6 5.5 ESTs H98153 RC_A4053886 RC_A4053886_s 1.2 5.5 ESTs; Highly similar to CYTOSOL AMINOPEPTIDASE [Bos	•				5.8
quinoid dihydropteridine reductase T69009 RC_T69009_s 0.5 5.8 ESTs ESTs Highly similar to follistatin-related protein [H.sapiens] D51110 RC_D51110_s 2.1 5.7 ESTs; Highly similar to follistatin-related protein [H.sapiens] D51110 RC_D51110_s 2.1 5.7 ESTs; Moderately similar to !!!! ALU SUBFAMILY SP WARNING ENTRY !!!! [H.sapiens] RC_AA452855 RC_AA452855 2.0 5.7 ESTs; Moderately similar to !!!! ALU SUBFAMILY SP WARNING ENTRY !!!! [H.sapiens] RA442125 RC_AA442125 RC_AA422125 RC_AA442125 RC_AA422125 RC_AA422125 RC_AA422125 RC_AA422125 RC_AA422125 RC_AA422125 RC_AA422125 RC_AA422125 RC_AA43222 RC_AA43222 RC_AA43222 RC_AA43222 RC_AA43222 RC_AA43222 RC_AA43222 RC_AA43222 RC_AA43222 RC_AA442123 RC_AA441223 RC_AA441124 RC_AA41124 RC_AA		T97257			5.8
ESTs					5.8
ESTs; Highly similar to follistatin-related protein [H.sapiens] D51110 RC_D51110_s 2.1 5.7   ESTs AA452855 RC_AA452855 2.0 5.7   ESTs Moderately similar to !!!! ALU SUBFAMILY SP WARNING ENTRY !!! [H.sapiens]   AA442125 RC_AA442125 1.3 5.7   Cell division cycle 42 (GTP-binding protein; 25kD) N63172 RC_N63172 2.1 5.7   ESTs; Highly similar to EUKARYOTIC INITIATION FACTOR 4 GAMMA [Onyclolagus cuniculus] W84870 RC_W84870_s 1.2 5.7   ESTS AA421121 RC_AA121121 1.3 5.7   ESTS; Moderately similar to HN1 [M.musculus] AA436027 RC_AA436027 1.9 5.7   ESTS; Weakly similar to brain-specific L-proline transporter AA460049 AA460049_s 1.2 5.7   ESTS; Weakly similar to brain-specific L-proline transporter AA460049 AA460049_s 1.2 5.7   ESTS AA453783 RC_AA453783_s 3.7 5.6   ESTS; Weakly similar to (defline not available 446577) [H.sapiens] AA486655 RC_AA486655 2.3 5.6   ESTS; Weakly similar to (defline not available 446577) [H.sapiens] AA496914 RC_AA496914 0.8 5.6   ESTS; Weakly similar to (defline not available 446577) [H.sapiens] AA496914 RC_AA496914 0.8 5.6   ESTS; Weakly similar to (defline not available 446577) [H.sapiens] AA496914 RC_AA496914 0.8 5.6   ESTS; Weakly similar to (defline not available 446577) [H.sapiens] AA496914 RC_AA496914 0.8 5.6   ESTS; Weakly similar to (defline not available 446577) [H.sapiens] AA496914 RC_AA496914 0.8 5.6   ESTS; Weakly similar to (defline not available 446577) [H.sapiens] AA496914 RC_AA496914 0.8 5.6   ESTS AA496914 RC_AA496914 0.8 5.6   ESTS Human mRNA for KIAA69 gene; partial cds AA48318 RC_AA48318 RC_AA48318 RC_AA48318 RC_AA48318 RC_AA483461 1.0   ESTS Hymportin alpha 1 (Importin alpha 5) N35247 RC_N35247 1.2 5.6   Homo sapiens clone 23675 mRNA sequence AA453461 RC_H73161 RC_H7		AA489459	<del>-</del>	1.1	5.7
ESTs	ESTs: Highly similar to follistatin-related protein [H.sapiens]	D51110	<del>-</del>	2.1	5.7
ESTs; Moderately similar to !!!! ALU SUBFAMILY SP WARNING ENTRY !!!! [H. saplens]		AA452855	<del>-</del> -	2.0	5.7
cell division cycle 42 (GTP-binding protein; 25kD)         N63172         RC_N63172         2.1         5.7           ESTs; Highly similar to EUKARYOTIC INITIATION FACTOR 4 GAMMA [Oryctolagus cuniculus]         W84870         RC_W84870_s         1.2         5.7           ESTs         AA121121         RC_AA121121         1.3         5.7           ESTs; Moderately similar to HN1 [M.musculus]         AA436027         RC_AA436027         1.9         5.7           ESTs         AA441923         RC_AA441923         5.6         5.7           ESTs         AA460049         AA460049_s         1.2         5.7           ESTs         AA4503962         RC_AA481911         0.7         5.6           ESTs         AA453783         RC_AA053962         1.2         5.6           ESTs         AA453783         RC_AA453783_s         3.7         5.6           ESTs         N32811         RC_N32811         1.8         5.6           ESTs         N32811         RC_N4386655         2.3         5.6           ESTs; Weakly similar to (defline not available 446577)         RC_AA486655         2.3         5.6           ESTs; Weakly similar to form transcription factor C-MAF (c-maf)         RC_AA496914         RC_AA496914         0.8         5.6	ESTs; Moderately similar to !!!! ALU SUBFAMILY SP WARNING		_	1.3	5.7
ESTs; Highly similar to EUKARYOTIC INITIATION FACTOR 4   GAMMA (Oryctolagus cuniculus)   W84870   RC_W84870_s   1.2 5.7   ESTs   AA121121   RC_AA121121   1.3 5.7   ESTs; Moderately similar to HN1 [M.musculus]   AA436027   RC_AA436027   1.9 5.7   ESTs   AA441923   RC_AA441923   RC_AA441923   RC_AA441923   RC_AA441923   RC_AA441923   RC_AA441923   RC_AA441923   RC_AA441923   RC_AA481911   RC_AA181911   RC	1	N63172	RC_N63172	2.1	5.7
ESTs	ESTs; Highly similar to EUKARYOTIC INITIATION FACTOR 4	1440.4070	_	4.0	
ESTs; Moderately similar to HN1 [M.musculus]					1
ESTs			<del>-</del>		- (
ESTs; Weakly similar to brain-specific L-proline transporter  AA460049  AA4603962  RC_AA053962  RC_AA053962  RC_AA453783  RC_AA453783  RC_AA453783  RC_AA453783  RC_AA453783  RC_AA453783  RC_AA485655  RC_AA453661  RC_AA453661  RC_AA453461  RC_AA453			<del>-</del>		1
ESTs	1		_		1
ESTs			<del>-</del>		1
ESTs	1		<del></del>		•
ESTs N32811 RC_N32811 1.8 5.6 Human transposon-like element mRNA M23161 M23161 0.8 5.6 ESTs AA485655 RC_AA485655 2.3 5.6 ESTs; Weakly similar to (defline not available 446577) [H.sapiens] Z41803 RC_Z41803 1.1 5.6 Homo sapiens short form transcription factor C-MAF (c-maf) mRNA; complete cds AA496914 RC_AA496914 0.8 5.6 glioblastoma amplified sequence AA095021 AA095021 1.0 5.6 karyopherin alpha 1 (importin alpha 5) N35247 RC_N35247 1.2 5.6 Human mRNA for KIAA69 gene; partial cds AA148318 RC_AA148318_s 2.3 5.6 Homo sapiens clone 23675 mRNA sequence Z39978 RC_Z39978 1.1 5.6 ESTs H73161 RC_H73161_f 1.2 5.6 Homo sapiens clone 23585 mRNA sequence AA453461 RC_AA453461 1.6 5.5 ESTs H98153 RC_H98153 7.0 5.5 desmoplakin (DPI; DPII) H90899 RC_H90899 5.4 5.5 sterol regulatory element binding transcription factor 2 AA053886 RC_AA053886_s 1.2 5.5			<del>-</del>		
Human transposon-like element mRNA M23161 M23161 0.8 5.6 ESTs AA485655 RC_AA485655 2.3 5.6 ESTs; Weakly similar to (defline not available 446577) [H.sapiens] Z41803 RC_Z41803 1.1 5.6 Homo sapiens short form transcription factor C-MAF (c-maf) mRNA; complete cds AA496914 RC_AA496914 0.8 5.6 glioblastoma amplified sequence AA095021 AA095021 1.0 5.6 karyopherin alpha 1 (importin alpha 5) N35247 RC_N35247 1.2 5.6 Human mRNA for KIAA69 gene; partial cds AA148318 RC_AA148318_s 2.3 5.6 Homo sapiens clone 23675 mRNA sequence Z39978 RC_Z39978 1.1 5.6 ESTs H73161 RC_H73161_f 1.2 5.6 Homo sapiens clone 23585 mRNA sequence AA453461 RC_AA453461 1.6 5.5 ESTs H98153 RC_H98153 7.0 5.5 desmoplakin (DPI; DPII) H90899 RC_H90899 5.4 5.5 sterol regulatory element binding transcription factor 2 AA053886 RC_AA053886_s 1.2 5.5	f				1
ESTs			<del>-</del>		
ESTs; Weakly similar to (defline not available 446577) [H.sapiens] Z41803 RC_Z41803 1.1 5.6 Homo sapiens short form transcription factor C-MAF (c-maf) mRNA; complete cds A4496914 RC_A4496914 0.8 5.6 glioblastoma amplified sequence AA095021 AA095021 1.0 5.6 karyopherin alpha 1 (importin alpha 5) N35247 RC_N35247 1.2 5.6 Human mRNA for KIAA69 gene; partial cds AA148318 RC_AA148318_s 2.3 5.6 Homo sapiens clone 23675 mRNA sequence Z39978 RC_Z39978 1.1 5.6 ESTs H73161 RC_H73161_f 1.2 5.6 Homo sapiens clone 23585 mRNA sequence AA453461 RC_A453461 1.6 5.5 ESTs H98153 RC_H98153 7.0 5.5 desmoplakin (DPI; DPII) H90899 RC_H90899 5.4 5.5 sterol regulatory element binding transcription factor 2 AA053886 RC_AA053886_s 1.2 5.5	· · · · · · · · · · · · · · · · · · ·				í
Homo sapiens short form transcription factor C-MAF (c-maf) mRNA; complete cds	ESTs; Weakly similar to (defline not available 446577)		_		5.6
glioblastoma amplified sequence       AA095021       AA095021       1.0       5.6         karyopherin alpha 1 (importin alpha 5)       N35247       RC_N35247       1.2       5.6         Human mRNA for KIAA69 gene; partial cds       AA148318       RC_AA148318_s       2.3       5.6         Homo sapiens clone 23675 mRNA sequence       Z39978       RC_Z39978       1.1       5.6         ESTs       H73161       RC_H73161_f       1.2       5.6         Homo sapiens clone 23585 mRNA sequence       AA453461       RC_AA453461       1.6       5.5         ESTs       H98153       RC_H98153       7.0       5.5         desmoplakin (DPI; DPII)       H90899       RC_H90899       5.4       5.5         sterol regulatory element binding transcription factor 2       AA053886       RC_AA053886_s       1.2       5.5         ESTs; Highly similar to CYTOSOL AMINOPEPTIDASE [Bos	Homo sapiens short form transcription factor C-MAF (c-maf)		_	•	l
karyopherin alpha 1 (importin alpha 5)       N35247       RC_N35247       1.2       5.6         Human mRNA for KIAA69 gene; partial cds       AA148318       RC_AA148318_s       2.3       5.6         Homo sapiens clone 23675 mRNA sequence       Z39978       RC_Z39978       1.1       5.6         ESTs       H73161       RC_H73161_f       1.2       5.6         Homo sapiens clone 23585 mRNA sequence       AA453461       RC_AA453461       1.6       5.5         ESTs       H98153       RC_H98153       7.0       5.5         desmoplakin (DPI; DPII)       H90899       RC_H90899       5.4       5.5         sterol regulatory element binding transcription factor 2       AA053886       RC_AA053886_s       1.2       5.5         ESTs; Highly similar to CYTOSOL AMINOPEPTIDASE [Bos			<del>-</del>		5.6
Human mRNA for KIAA69 gene; partial cds       AA148318       RC_AA148318_s       2.3       5.6         Homo sapiens clone 23675 mRNA sequence       Z39978       RC_Z39978       1.1       5.6         ESTs       H73161       RC_H73161_f       1.2       5.6         Homo sapiens clone 23585 mRNA sequence       AA453461       RC_AA453461       1.6       5.5         ESTs       H98153       RC_H98153       7.0       5.5         desmoplakin (DPI; DPII)       H90899       RC_H90899       5.4       5.5         sterol regulatory element binding transcription factor 2       AA053886       RC_AA053886_s       1.2       5.5         ESTs; Highly similar to CYTOSOL AMINOPEPTIDASE [Bos	1				ł
Homo sapiens clone 23675 mRNA sequence         Z39978         RC_Z39978         1.1         5.6           ESTs         H73161         RC_H73161_f         1.2         5.6           Homo sapiens clone 23585 mRNA sequence         AA453461         RC_AA453461         1.6         5.5           ESTs         H98153         RC_H98153         7.0         5.5           desmoplakin (DPI; DPII)         H90899         RC_H90899         5.4         5.5           sterol regulatory element binding transcription factor 2         AA053886         RC_AA053886_s         1.2         5.5           ESTs; Highly similar to CYTOSOL AMINOPEPTIDASE [Bos					- 1
ESTs         H73161         RC_H73161_f         1.2         5.6           Homo sapiens clone 23585 mRNA sequence         AA453461         RC_AA453461         1.6         5.5           ESTs         H98153         RC_H98153         7.0         5.5           desmoplakin (DPI; DPII)         H90899         RC_H90899         5.4         5.5           sterol regulatory element binding transcription factor 2         AA053886         RC_AA053886_s         1.2         5.5           ESTs; Highly similar to CYTOSOL AMINOPEPTIDASE [Bos					i i
Homo sapiens clone 23585 mRNA sequence         AA453461         RC_AA453461         1.6         5.5           ESTs         H98153         RC_H98153         7.0         5.5           desmoplakin (DPI; DPII)         H90899         RC_H90899         5.4         5.5           sterol regulatory element binding transcription factor 2         AA053886         RC_AA053886_s         1.2         5.5           ESTs; Highly similar to CYTOSOL AMINOPEPTIDASE [Bos         Bos         Bos         Bos         Bos         Bos	,		<del>-</del>		5.6
ESTs         H98153         RC_H98153         7.0         5.5           desmoplakin (DPI; DPII)         H90899         RC_H90899         5.4         5.5           sterol regulatory element binding transcription factor 2         AA053886         RC_AA053886_s         1.2         5.5           ESTs; Highly similar to CYTOSOL AMINOPEPTIDASE [Bos         5.5         5.5         5.5         5.5			<del>-</del>		5.5
desmoplakin (DPI; DPII) H90899 RC_H90899 5.4 5.5 sterol regulatory element binding transcription factor 2 AA053886 RC_AA053886_s 1.2 5.5 ESTs; Highly similar to CYTOSOL AMINOPEPTIDASE [Bos	•				5.5
sterol regulatory element binding transcription factor 2 AA053886 RC_AA053886_s 1.2 5.5 ESTs; Highly similar to CYTOSOL AMINOPEPTIDASE [Bos			<del>-</del>		5.5
			<del>-</del>		5.5
		AA134138	RC_AA134138	1.4	5.5

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ESTs; Highly similar to PROBABLE UBIQUITIN CARBOXYL-TERMINAL HYDROLASE [Mus musculus]	AA082057	RC_AA082057	1.1	5.5
ESTs; Highly similar to HYPOTHETICAL GTP-BINDING PROTEIN IN PMI4-PAC2 INTERGENIC REGION				
[Saccharomyces cerevisiae]	AA609710	RC_AA609710	5.5	5.5
heterogeneous nuclear ribonucleoprotein A1	AA416785	RC_AA416785_f	2.2	5.5
Human Chromosome 16 BAC clone CIT987SK-A-362G6	U95740	U95740_rna1	0.9	5.5
Homo sapiens Arp2/3 protein complex subunit p21-Arc (ARC21) mRNA; complete cds	AF006086	AF006086	1.4	5.4
ESTs	AA478387	RC_AA478387	1.0	5.4
Homo sapiens multiple membrane spanning receptor TRC8 (TRC8) mRNA; complete cds	AA455970	RC_AA455970	1.8	5.4
ESTs; Weakly similar to NIPSNAP2 protein [H.sapiens]	R49052	RC_R49052	-1.3	5.4
ESTs	AA235803	RC_AA235803_i	2.5	5.4
ESTs	T15482	RC_T15482_f	0.7	5.4
NADH dehydrogenase (ubiquinone) 1 beta subcomplex; 3 (12kD; B12)	AA040759	RC_AA040759_s	1.0	5.4
Human mRNA for KIAA263 gene; complete cds	T16989	RC_T16989_f	1.2	5.4
ESTs	R27975	RC_R27975	1.2	5.4
ESTs; Moderately similar to (defline not available 445515)		_		
[H.sapiens]	AA047187	RC_AA047187	8.0	5.4
ESTs	W45417	RC_W45417	1.0	5.4
ESTs; Weakly similar to zinc finger protein [H.sapiens]	AA487297	RC_AA487297	1.6	5.4
	M27830	AFFX-M27830_5	0.6	5.4
ESTs	AA258614	RC_AA258614_s	2.0	5.3
ESTs; Weakly similar to cDNA EST EMBL:T1157 comes from this gene [C.elegans]	AA313414	AA313414_s	1.5	5.3
secreted frizzled-related protein 4	AA291725	RC_AA291725	5.3	5.3
ESTs	AA282179	RC_AA282179	0.9	5.3
Human pim-2 protooncogene homolog pim-2h mRNA; complete	AA227480	RC_AA227480_s	0.8	5.3
ESTs; Weakly similar to ORF YOR126c [S.cerevisiae]	AA249311	AA249311	1.4	5.3
ESTs	W36290	W36290_s	1.7	5.3
Homo sapiens hJTB mRNA; complete cds	AA071387	AA071387	1.7	5.3
Homo sapiens mRNA for putative vacuolar proton ATPase membrane sector associated protein M8-9	D51241	RC_D51241_s	2.2	5.3
ESTs	T92735	RC_T92735	1.7	5.3
splicing factor proline/glutamine rich (polypyrimidine tract-binding		_		
protein-associated)	T64923	RC_T64923_f	1.2	5.3
ESTs	W20404	RC_W20404_s	1.1	5.2
high-mobility group (nonhistone chromosomal) protein 1	AA486201	RC_AA486201_s	1.1	5.2
ESTs	T15386	RC_T15386	0.9	5.2
ESTs	AA608657	RC_AA608657_f	2.1	5.2
ESTs	AA236276	RC_AA236276	1.4	5.2
ESTs; Weakly similar to cDNA EST EMBL:T1585 comes from this gene [C.elegans]	AA598439	RC_AA598439	1.4	5.2
ubiquitin-specific protease 1	D62657	RC_D62657	0.5	5.2
ESTs	AA419507	AA419507	1.0	5.2
ESTs	AA496962	RC_AA496962	0.9	5.2
Homo sapiens clone 23596 mRNA sequence	AA425741	RC_AA425741	0.7	5.2
Homo sapiens clone 23714 mRNA sequence	AA147364	RC_AA147364	0.9	5.2
ESTs; Moderately similar to !!!! ALU SUBFAMILY J WARNING ENTRY !!!! [H.sapiens]	Z21420	Z21420	1.3	5.2

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ESTs	T15434	RC_T15434_s	0.9	5.1
ESTs	W95416	RC_W95416	1.1	5.1
Homo sapiens clone 23675 mRNA sequence	AA018804	AA018804	2.9	5.1
Homo sapiens mRNA for KIAA447 protein; complete cds	C02016	C02016	1.4	5.1
apolipoprotein H (beta-2-glycoprotein I)	T83356	RC_T83356_s	0.3	5.1
ESTs; Moderately similar to weak similarity to Arabidopsis thaliana ubiquitin-like protein 8 [C.elegans]	D31544	D31544_s	3.2	5.1
Homo sapiens MAD-related gene SMAD7 (SMAD7) mRNA; complete cds	AF010193	AF010193	1.0	5.1
ESTs; Weakly similar to !!!! ALU SUBFAMILY J WARNING ENTRY !!!! [H.sapiens]	AA155779	RC_AA155779	1.7	5.1
ESTs; Highly similar to 6S RIBOSOMAL PROTEIN L26 [Homo sapiens; Mus musculus]	D80128	RC_D80128_f	1.6	5.1
ESTs	AA018907	RC_AA018907_s	2.0	5.1
immunoglobulin gamma 3 (Gm marker)	J00231	J00231_f	1.4	5.1
ESTs	N81162	N81162	2.0	5.1
ESTs	AA599850	RC_AA599850	1.3	5.1
ESTs	AA460935	RC_AA460935	1.8	5.0
ESTs; Highly similar to heat shock factor binding protein 1 HSBP1 [H.sapiens]	AA490864	RC_AA490864	1.4	5.0
ESTs	W80516	W80516	1.0	5.0
ESTs; Weakly similar to neuronal thread protein AD7c-NTP [H.sapiens]	AA046939	RC_AA046939_s	1.2	5.0
Human DNA sequence from clone 149A16 on chromosome 22q12-13. Contains an IGLC (Immunoglobulin Lambda Chain C) pseudogene; the RFPL3 and RFPL3S genes for Ret finger protein-like 3 and Ret finger protein-like 3 antisense respectively a gene for a novel Imm		RC_AA151882	,1.4	5.0
ESTS	T72867	RC_T72867	1.2	5.0
myosin VI	AB002387	AB002387	4.5	5.0
GM2 ganglioside activator protein	AA167512	RC_AA167512	1.3	5.0
ESTs; Moderately similar to putative p15 [H.sapiens]	AA481060	RC_AA481060	1.3	5.0
ESTs	N69086	RC_N69086	1.5	5.0
ESTs; Highly similar to heat shock factor binding protein 1	04.40.40	DC C14243 f	1.7	5.0
HSBP1 [H.sapiens]	C14243	RC_C14243_f	1.4	5.0
neuroblastoma RAS viral (v-ras) oncogene homolog	AA431977	RC_AA431977	1.4	5.0
Homo sapiens mRNA for putative vacuolar proton ATPase membrane sector associated protein M8-9	R25326	R25326	0.9	5.0
butyrate response factor 1 (EGF-response factor 1)	H40424	RC_H40424_s	1.4	5.0
H factor (complement)-like 1	AA235873	RC_AA235873_s	0.6	5.0
ESTs; Weakly similar to predicted using Genefinder [C.elegans]	AA252040	RC_AA252040	1.5	5.0
ESTs	R62589	RC_R62589_f	1.2	5.0
Human mRNA for KIAA171 gene; complete cds	AA028889	RC_AA028889_s	1.1	5.0
ESTs; Highly similar to G protein-coupled receptor kinase 6; splice variant B [H.sapiens]	AA040699	RC_AA040699	1.0	4.9
ESTs; Highly similar to (defline not available 45813) [H.sapiens]		RC_AA488414	1.2	4.9
CYTOCHROME C	M22877	M22877	0.8	4.9
ESTs; Highly similar to synapsin I [R.norvegicus]	T15663	RC_T15663_s	0.8	4.9
lysozyme (renal amyloidosis)	J03801	J03801_f	0.9	4.9
ESTs	202001			
[E318	W88642	RC_W88642	1.0	4.9
		RC_W88642 RC_W57813_i	1.0 0.8	4.9 4.9
ESTs ESTs	W88642			

#### FIGURE 8

(Cont.)

Homo sapiens mRNA for KIAA829 protein; partial cds	H01766	H01766_s	1.3	4.9
ESTs; Weakly similar to synapse associated protein sap47-2 [D.melanogaster]	N51752	RC_N51752	2.5	4.9
secreted phosphoprotein 1 (osteopontin; bone sialoprotein I;	1120759	_	4.0	
early T-lymphocyte activation 1)	U20758 AA283085	U20758_rna1 RC_AA283085 s	1.9	4.9
ESTs			0.6	4.9
ATPase; Ca++ transporting; cardiac muscle; slow twitch 2	M23114 AA600140	M23114	2.0	4.9
Deleted in oral cancer-1	W88755	RC_AA600140	2.4	4.9
ESTs	C13990	RC_W88755	1.3	4.8
synuclein; alpha (non A4 component of amyloid precursor)	R73982	RC_C13990_f R73982	0.6 0.7	4.8
ESTs; Weakly similar to KIAA638 protein [H.sapiens] ESTs	AA281949	RC_AA281949		4.8 4.8
ESTs	AA281949 AA134767	<del>-</del>	1.3	4.0 4.8
ESTs	AA453593	RC_AA134767_s	1.5 0.9	
frizzled (Drosophila) homolog 1	AA358618	RC_AA453593_s RC_AA358618_s	2.4	4.8 4.8
ESTs	H93552	_		4.8
	D53233	RC_H93552 RC_D53233	0.9 2.7	4.8
cyclin G2 peptidylprolyl isomerase B (cyclophilin B)	H15847	RC_D53233 RC_H15847_s	1.8	4.8
ESTs; Highly similar to (defline not available 439889)	H15047	KC_H15647_S	1.0	4.0
[H.sapiens]	AA256210	RC_AA256210	2.0	4.8
Homo sapiens clone 23698 mRNA sequence	R32440	RC_R32440	0.5	4.8
ESTs	C15078	RC_C15078_i	0.8	4.8
ESTs; Weakly similar to CH-TOG PROTEIN [H.sapiens]	AA486092	RC_AA486092	2.8	4.8
ESTs	C00038	 C00038_s	2.8	4.8
ESTs	AA450281	RC_AA450281	1.0	4.8
Homo sapiens mRNA for KIAA663 protein; complete cds	N57577	RC_N57577	1.1	4.8
ferritin; light polypeptide	T73572	RC_T73572_f	1.1	4.8
ESTs; Highly similar to 5'-AMP-ACTIVATED PROTEIN KINASE; GAMMA-1 SUBUNIT [Rattus norvegicus]	AA114970	RC_AA114970_i	1.3	4.8
ESTs; Highly similar to ZYXIN [Gallus gallus]	H25769	RC_H25769_s	1.2	4.8
tubulin-specific chaperone a	AA504095	AA504095	1.6	4.8
ESTs	N25576	RC_N25576	1.1	4.8
Homo sapiens inner mitochondrial membrane translocase Tim23 (TIM23) mRNA; nuclear gene encoding mitochondrial protein;	}	_		
complete cds	AA442768	RC_AA442768_i	1.9	4.8
ESTs	AA235289	RC_AA235289	2.4	4.8
ESTs	N27198	RC_N27198	2.5	4.8
ESTs	AA479132	AA479132	1.7	4.8
ESTs; Moderately similar to neuronal thread protein AD7c-NTP [H.sapiens]	AA450228	RC_AA450228	1.5	4.7
calmodulin 1 (phosphorylase kinase; delta)	AA085590	RC_AA085590_s	1.3	4.7
Homo sapiens actin-related protein Arp3 (ARP3) mRNA; complete cds	AA199588	RC_AA199588	1.8	4.7
Human TAR DNA-binding protein-43 mRNA; complete cds	H16390		1.3	4.7
Human TAR RNA loop binding protein (TRP-185) mRNA; complete cds	N70678	 RC_N70678_s	1.5	4.7
Human transformer-2 alpha (htra-2 alpha) mRNA; complete cds	AA455812	AA455812	1.3	4.7
ESTs	AA057287	AA057287	0.7	4.7
ESTs; Weakly similar to !!!! ALU SUBFAMILY SQ WARNING ENTRY !!!! [H.sapiens]	AA070801	RC_AA070801	6.3	4.7
ESTs; Highly similar to COMPLEMENT C1Q SUBCOMPONENT; A CHAIN PRECURSOR [Homo sapiens]	W87494	RC_W87494	1.2	4.7

Isot.	AA490264	PC AA400364	0.8	4.7
ESTs	AA303745	RC_AA490264 AA303745_s		4.7
TAP binding protein (tapasin)	AA454086	***	1.8	4.7
UDP-glucose dehydrogenase	Y00062	RC_AA454086_f Y00062	1.3	4.7
protein tyrosine phosphatase; receptor type; c polypeptide			0.6	4.6
ESTS	AA449333	RC_AA449333	2.9	4.6
ESTs; Highly similar to TURNED ON AFTER DIVISION; 64 KD PROTEIN [Rattus norvegicus]	AA058664	RC_AA058664	1.5	4.6
ESTs	R51913	RC_R51913	1.3	4.6
protein tyrosine phosphatase type IVA; member 2	AA329274	AA329274_f	1.9	4.6
calumenin	W84712	RC_W84712	3.5	4.6
ESTs	AA173597	AA173597	1.8	4.6
H.sapiens mRNA for Fas/Apo-1 (clone pCRTM11-Fasdelta(4,7))	X83492	X83492	0.8	4.6
ESTs	AA404427	RC_AA404427	1.3	4.6
ESTs	W65477	RC_W65477	1.2	4.6
ESTs	AA040270	RC_AA040270	1.3	4.6
hemoglobin; gamma A	R92458	RC_R92458_f	0.5	4.6
ESTs	F01831	RC_F01831	0.2	4.6
ESTs	Z41372	RC_Z41372_s	2.0	4.6
ESTs; Weakly similar to PROBABLE ATP-DEPENDENT RNA HELICASE HRH1 [H.sapiens]	AA410336	RC_AA410336	2.0	4.6
ESTs; Highly similar to GENERAL NEGATIVE REGULATOR OF TRANSCRIPTION SUBUNIT 2 [Saccharomyces cerevisiae]	AA010686	AA010686	1.6	4.6
ESTs; Weakly similar to keratin 1 [H.sapiens]	AA037386	RC_AA037386_s	1.1	4.6
H1 histone family; member 2	T90190	RC_T90190_s	1.5	4.5
lysozyme (renal amyloidosis)	X14008	X14008_rna1_f	0.9	4.5
ESTs	Z39622	RC_Z39622_s	2.1	4.5
ribosomal protein L22	T23926	RC_T23926	1.5	4.5
tumor necrosis factor (ligand) superfamily; member 1	H25836	RC_H25836	3.2	4.5
ESTs	R60952	RC_R60952_i	1.0	4.5
Homo sapiens clone 23836 mRNA sequence	T17428	RC_T17428_s	1.0	4.5
NADH dehydrogenase (ubiquinone) 1 beta subcomplex; 7 (18kD; B18)	AA609299	RC_AA609299_s	1.1	4.5
SET PROTEIN	AA205665	RC_AA205665_s	1.7	4.5
Human mRNA for KIAA349 gene; partial cds	AB002347	AB002347	0.7	4.5
protease; serine; 11 (IGF binding)	T82292	RC_T82292_s	2.1	4.5
ESTs	AA465218	RC_AA465218	1.6	4.5
ESTs	AA236018	RC_AA236018	1.2	4.5
SRY (sex determining region Y)-box 4	AA479953	RC_AA479953	8.7	4.5
LIVER CARBOXYLESTERASE PRECURSOR	T68878	RC_T68878_f	0.3	4.4
Human mRNA for KIAA228 gene; partial cds	AA431206	RC_AA431206_s	1.5	4.4
ESTs	AA489012	RC_AA489012	2.4	4.4
ESTs; Highly similar to NUCLEAR FACTOR 1 A1 [Gallus gallus]	F08945	RC_F08945	1.5	4.4
zinc finger protein 27	H80409	RC_H80409	1.5	4.4
ESTs	T86337	RC_T86337	1.2	4.4
ESTs	AA459245	RC_AA459245	0.9	4.4
ribosomal protein L22	N93380	RC_N93380	1.3	4.4
ATPase; Ca++ transporting; cardiac muscle; slow twitch 2	W61297	RC_W61297	0.4	4.4
ESTs	AA291749	RC_AA291749_s	4.4	4.4
tubulin; beta polypeptide	T03651	RC_T03651_s	1.0	4.4
regulator of G-protein signalling 5	AA348466	RC_AA348466_s	1.4	4.4

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ESTs; Highly similar to ARGINYL-TRNA SYNTHETASE; MITOCHONDRIAL PRECURSOR [Saccharomyces cerevisiae]	R12777	RC_R12777_s	1 =	ارر
cytochrome c oxidase subunit VIIb	Z14244	Z14244	1.5 0.9	4.4
ESTs	T92718	RC_T92718	1.1	4.4
protein phosphatase 2; regulatory subunit B (B56); delta isoform	T96379	RC_T96379_s	1.4	4.3
LYSOSOME-ASSOCIATED MEMBRANE GLYCOPROTEIN 1	100075	1/0_130379_5	1.4	4.3
PRECURSOR	T74571	RC_T74571_s	1.2	4.3
immunoglobulin lambda-like polypeptide 2	M34516	M34516_r	1.1	4.3
ESTs; Moderately similar to unknown [H.sapiens]	N23222	RC_N23222	2.2	4.3
ESTs	Z38688	RC_Z38688	0.3	4.3
ESTs	W42412	RC_W42412	1.1	4.3
ESTS; Highly similar to HEMOPOIETIC-SPECIFIC EARLY	1470407	DO 1400407 -		
RESPONSE PROTEIN [Mus musculus]	W20487	RC_W20487_s	1.4	4.3
ESTs	N21407	RC_N21407	1.8	4.3
ESTs; Weakly similar to deduced amino acid sequence is highly homologous to hypothetical proteins of C.elegans(T23g5.4 and				
T23G5.2). [H.sapiens]	H97159	RC_H97159	1.2	4.3
poly (ADP-ribose) glycohydrolase	R69293	RC_R69293	1.0	4.3
ESTs	AA253459	RC_AA253459	0.9	4.3
ESTs	AA452248	RC_AA452248	0.9	4.3
Homo sapiens clone 23742 mRNA; partial cds	AA608649	RC_AA608649	1.2	4.3
ESTs; Highly similar to (defline not available 446693)				
[H.sapiens]	AA393432	AA393432_s	1.5	4.3
ESTs	AA398318	RC_AA398318	1.5	4.3
H.sapiens mRNA for translin associated zinc finger protein-1	R79723	RC_R79723_s	1.5	4.3
ESTs	Z38874	RC_Z38874	1.1	4.3
testis enhanced gene transcript	AA079500	RC_AA079500	1.1	4.3
ESTs	N48000	RC_N48000	2.7	4.3
von Hippel-Lindau syndrome ESTs	W31600 AA156230	RC_W31600_f	2.3	4.3
v-Ki-ras2 Kirsten rat sarcoma 2 viral oncogene homolog	H69138	RC_AA156230 RC_H69138	1.0 0.9	4.3
UDP-N-acetyl-alpha-D-galactosamine:polypeptide	1109130	VC_109138	0.9	4.3
N-acetylgalactosaminyltransferase 1 (GalNAc-T1)	T34527	T34527	2.6	4.3
ESTs; Highly similar to ZINC FINGER PROTEIN 91 [Homo				
sapiens]	AA029288	RC_AA029288	1.5	4.2
ESTs ESTs	AA458959	RC_AA458959	1.7	4.2
ESTs	T10108	RC_T10108_s	1.4	4.2
ras homolog gene family; member H	H04753	RC_H04753_f	3.2	4.2
Homo sapiens voltage dependent anion channel protein mRNA;	W63747	RC_W63747	1.8	4.2
complete cds	AA094989	AA094989	0.7	4.2
ESTs; Weakly similar to Bat2 [H.sapiens]	H05631	RC_H05631_f	1.2	4.2
ESTs	R44357	RC_R44357	1.8	4.2
ESTs	N34482	RC_N34482	1.4	4.2
Microfibril-associated glycoprotein-2	U37283	U37283 ·	1.1	4.2
ESTs; Weakly similar to CH-TOG PROTEIN [H.sapiens]	AA418985	RC_AA418985	2.3	4.2
ESTs; Moderately similar to (defline not available 4589678)	4.4050705	DO 44050707	4.6	
[H.sapiens]	AA252765	RC_AA252765	1.0	4.2
Homo sapiens mRNA for KIAA214 protein; complete cds zinc finger protein 262	R24483	RC_R24483_s	0.6	4.2
CAAX box 1	AA481428	RC_AA481428	1.0	4.2
TOLOGO POY I	AA279811	RC_AA279811_s	0.9	4.2

ESTs; Weakly similar to (defline not available 46333)	AA452082	A A 462002	4.0	
[H.sapiens]		AA452082	1.0	4.
cytochrome b-561	T03441	RC_T03441_f	1.2	4.
ESTs	AA040945	RC_AA040945	0.8	4.
FSHD region gene 1	H82532	RC_H82532	1.1	4.
ESTs	AA093977	AA093977	1.2	4.
ESTs	R26589	RC_R26589_f	1.0	4.
von Hippel-Lindau binding protein 1	U56833	U56833	1.4	4.
S1 calcium-binding protein A8 (calgranulin A)	M21005	M21005	0.9	4.
ESTs; Highly similar to POLYADENYLATE-BINDING PROTE [Homo sapiens]	=IN W95592	RC_W95592_i	1.3	4.
ESTs	AA620962	RC_AA620962	1.2	4.
histatin 1	L05512	L05512	0.8	4.
ESTs	AA293426	RC_AA293426	0.7	4.
cyclin F	T89627	RC_T89627_s	1.3	4.
ESTs; Weakly similar to Similarity to Serpentwood strictosidin				
synthase precursor [C.elegans]	AA256171	RC_AA256171	1.9	4.
ESTs	AA311352	AA311352_s	1.6	4.
ESTs	AA405654	RC_AA405654_s	1.5	4.
ESTs; Highly similar to ACTIN II [Plasmodium falciparum]	AA040263	RC_AA040263	1.1	4.
ESTs	C01552	C01552	1.0	4.
ESTs; Weakly similar to KIAA319 [H.sapiens]	N95507	N95507	1.3	4.
ESTs	N63706	RC_N63706	1.0	4
STs	AA152312	RC_AA152312	1.1	4
aminin; beta 1	M61916	M61916	1.5	4
ESTs	AA063431	RC_AA063431_f	0.8	4
ESTs	N39016	RC_N39016	1.3	4
ESTs; Moderately similar to neuronal thread protein AD7c-NT				_
[H.sapiens]	W45457	RC_W45457	1.2	4.
ESTs	C15324	RC_C15324_f	4.2	4.
ESTs	N46086	N46086_s	1.6	4.
ESTs	W45494	RC_W45494	1.0	4.
bone morphogenetic protein 6	AA598702	RC_AA598702	1.6	4.
cadherin 11 (OB-cadherin; osteoblast)	D21254	D21254_s	3.2	4.
ESTs; Weakly similar to ribokinase [E.coli]	T69020	RC_T69020_s	0.9	4.
ESTs; Moderately similar to !!!! ALU SUBFAMILY SX WARNI ENTRY !!!! [H.sapiens]	NG AA120783	RC_AA120783	1.4	4.
erythrocyte membrane protein band 7.2 (stomatin)	H27442	RC_H27442_s	1.0	4.
solute carrier family 7 (cationic amino acid transporter; y+				
system); member 6	R51116	RC_R51116_f	0.8	4.
ESTs	T69728	RC_T69728	1.1	4.
Homo sapiens clone 2377 mRNA sequence	R44163	RC_R44163_f	0.9	4.
ESTs	W80739	RC_W80739_f	1.0	4.
ESTs; Moderately similar to LNXp7 [M.musculus]	H82424	RC_H82424	1.7	4.
secreted frizzled-related protein 4	AA487193	RC_AA487193	4.7	4.
ESTs; Moderately similar to neuronal thread protein AD7c-NT [H.sapiens]	P AA428607	RC_AA428607	1.3	4.
fumarylacetoacetate	W79422	 RC_W79422_s	1.3	4.
MATRIN 3	N20178	RC_N20178	1.2	4.
ESTs	W67727	RC_W67727	1.4	4.
ESTs	AA011510	RC_AA011510	1.8	4.

# FIGURE 8 (Cont.)

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ESTs	AA069569	RC_AA069569	1.5	4.0
	N92934	RC_N92934_s	2.5	4.0
cysteine-rich protein 1 (intestinal) ESTs	AA004415	RC_AA004415	1.2	4.0
	W81205	<del>-</del>		4.0
ESTs	N56993	RC_W81205	1.5	Į.
ESTs		RC_N56993	2.0	4.0
ESTs	AA256943	RC_AA256943_s	0.8	4.0
ESTs	N68133	RC_N68133	0.7	4.0
homogentisate 1;2-dioxygenase (homogentisate oxidase)	R08615	RC_R08615_s	0.5	4.0
Meis1 (mouse) homolog	N95243	RC_N95243_s	0.9	4.0
Accession not listed in Genbank	K01160	K01160	1.7	4.0
ESTs	T33489	RC_T33489_s	1.3	4.0
H2A histone family; member Z	M37583	M37583	2.8	4.0
B-factor; properdin	T72268	RC_T72268_s	1.3	4.0
ESTs	AA133457	RC_AA133457	1.2	4.0
ESTs	AA287681	RC_AA287681_s	1.3	4.0
ESTs; Highly similar to HYPOTHETICAL 84.7 KD PROTEIN ZK198.1 IN CHROMOSOME III [Caenorhabditis elegans]	AA481403	RC_AA481403	4.0	4.0
ESTs	AA233445	RC_AA233445	1.9	4.0
ESTs; Weakly similar to PRE-MRNA SPLICING FACTOR		_		
SRP75 [Homo sapiens] ESTs; Weakly similar to deduced amino acid sequence is highly	AA452256	RC_AA452256	1.2	4.0
homologous to hypothetical proteins of C.elegans(T23g5.4 and				1
T23G5.2). [H.sapiens]	AA488433	RC_AA488433	1.1	4.0
Homo sapiens mRNA for KIAA719 protein; complete cds	R60689	RC_R60689	1.9	4.0
ESTs	AA016306	RC_AA016306	0.6	4.0
H.sapiens mRNA for nuclear protein SA-2	AA489057	RC_AA489057	6.2	4.0
ESTs	AA431191	RC_AA431191_s	1.8	4.0
protein kinase C; zeta	R24258	RC_R24258_s	0.7	4.0
ESTs	N66847	RC_N66847	1.4	4.0
ESTs	AA233548	RC_AA233548	1.5	4.0
ESTs	AA400229	RC_AA400229	1.7	4.0
ESTs; Weakly similar to SAS [H.sapiens]	H07011	H07011	1.8	3.9
Human mRNA for KIAA96 gene; partial cds	D60769	RC_D60769_s	0.9	3.9
cell division cycle 42 (GTP-binding protein; 25kD)	AA031548	AA031548	3.1	3.9
ESTs	H29293	RC_H29293_f	1.6	3.9
Rho GTPase activating protein 1	AA032067	RC_AA032067_s	2.0	3.9
ESTs; Highly similar to NADH-UBIQUINONE OXIDOREDUCTASE SUBUNIT B14.5B [Bos taurus]	AA234533	AA234533	1.4	3.9
calpain; large polypeptide L2	R39610	RC_R39610_s	1.3	3.9
ESTs	AA456845	RC_AA456845	1.4	3.9
ESTs; Highly similar to ATP SYNTHASE EPSILON CHAIN;	PV440040	RO_AA436645	1.4	3.9
MITOCHONDRIAL PRECURSOR [Bos taurus]	W72685	RC_W72685	1.3	3.9
Homo sapiens mRNA for KIAA886 protein; complete cds	W58081	RC_W58081	1.0	3.9
ESTs; Highly similar to (defline not available 467918) [H.sapiens]	AA026962	RC_AA026962	1.4	3.9
Human DNA from overlapping chromosome 19 cosmids R31396; F25451; and R3176 containing COX6B and UPKA;	T16050	DO T45050 £	2.0	2.0
genomic sequence	T15852	RC_T15852_f	2.0	3.9
ESTs	AA256317	RC_AA256317	1.3	3.9
ESTs	AA504492	RC_AA504492	2.4	3.9
ESTs	R78224	RC_R78224	1.0	3.9
ESTs	W74728	RC_W74728	0.2	3.9

# FIGURE 8 (Cont.)

<b>!</b>				
Human mRNA for KIAA263 gene; complete cds	T90946	RC_T90946_f	1.1	3.9
ESTs	D59711	RC_D59711_f	2.6	3.9
ESTs; Weakly similar to neuronal thread protein AD7c-NTP [H.sapiens]	AA428364	RC_AA428364_s	1.3	3.9
ESTs	AA342457	RC AA342457 i	2.1	3.9
ATPase; H+ transporting; lysosomal (vacuolar proton pump) 9kD		AA214710	1.4	3.9
	AA147532	RC_AA147532_s	2.8	3.9
Homo sapiens mRNA for nuclear protein; NP22; complete cds	AA157623	AA157623 s		
Homo sapiens mRNA for KIAA75 protein; complete cds		<del>-</del>	1.2	3.9
ESTs	T90345	RC_T90345	1.2	3.9
ESTs	AA410424	RC_AA410424	1.3	3.9
ESTs; Highly similar to (defline not available 467914) [H.sapiens]	N26691	RC_N26691	1.6	3.9
Homo sapiens mRNA for KIAA99 protein; partial cds	N51651	RC_N51651	0.9	3.9
ESTs; Highly similar to MICROSOMAL SIGNAL PEPTIDASE 21 KD SUBUNIT [Canis familiaris]	AA234347	RC_AA234347	1.3	3.9
ATPase; H+ transporting; lysosomal (vacuolar proton pump); beta polypeptide; 56/58kD; isoform 2	M60346	M60346_s	0.9	3.9
	X82554	X82554_rna1	0.8	3.9
RAB4; member RAS oncogene family ESTs; Highly similar to TISSUE ALPHA-L-FUCOSIDASE	X02334	A02334_Ma1	0.0	3.5
PRECURSOR [Homo sapiens]	AA234925	RC_AA234925	1.4	3.9
heterogeneous nuclear ribonucleoprotein A2/B1	AA131165	RC_AA131165_s	2.0	3.9
Human mariner-like element-containing mRNA; clone pcHMT1	AA487508	RC_AA487508	1.9	3.9
ESTs	AA489618	RC_AA489618_s	1.5	3.9
ESTs	AA436158	RC_AA436158	3.9	3.9
ESTs	AA256688	RC_AA256688_s	1.0	3.8
ESTs	H14982	RC_H14982_r	1.0	3.8
ADP-ribosyltransferase (NAD+; poly (ADP-ribose) polymerase)	J03473	J03473	2.1	3.8
ESTs	W60310	RC_W60310	1.3	3.8
ESTs	AA040397	RC_AA040397	1.5	3.8
ESTs	T15457	RC_T15457_f	1.0	3.8
ESTs; Weakly similar to neuronal thread protein AD7c-NTP [H.sapiens]	N67343	RC_N67343	2.1	3.8
ESTs; Highly similar to HYPOTHETICAL 3.5 KD PROTEIN				
C3A5.3 IN CHROMOSOME III [Caenorhabditis elegans]	AA348925	RC_AA348925_s	2.4	3.8
protocadherin 2 (cadherin-like 2)	T65540	RC_T65540_s	1.0	3.8
ESTs	AA404421	RC_AA404421	1.2	3.8
ESTs	AA237009	RC_AA237009	1.3	3.8
ESTs; Moderately similar to ATP-CITRATE [Rattus norvegicus]	D51405	RC_D51405	1.4	3.8
Human Ig J chain gene	M12759	M12759	0.5	3.8
ESTs; Moderately similar to alternatively spliced product using exon 13A [H.sapiens]	H90314	RC_H90314_s	1.2	3.8
NADH dehydrogenase (ubiquinone) Fe-S protein 4 (18kD) (NADH-coenzyme Q reductase)	AA063581	RC_AA063581	0.8	3.8
H.sapiens OZF mRNA	T25747	RC_T25747_s	1.8	3.8
lumican	U21128	U21128	2.2	3.8
heterogeneous nuclear ribonucleoprotein G	AA173143	RC_AA173143_s	1.6	3.8
ESTs	N63165	RC_N63165	1.1	3.8
ESTs; Weakly similar to KIAA62 [H.sapiens]	AA233763	RC_AA233763	0.6	3.8
Human high density lipoprotein binding protein (HBP) mRNA; complete cds	H28100	- RC_H28100_s	1.9	3.8
ESTs	AA074350	RC_AA074350	1.3	3.8
ESTs	W46632	RC_W46632	1.1	3.8
JE013	VV4003Z	NO_VV+003Z	1.1	3.0

# FIGURE 8 (Cont.)

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No.	EST\$	AA491278	RC_AA491278_r	1.0	3.8
AZ251902	ESTs; Highly similar to GASTRULA ZINC FINGER PROTEIN	TE0752	DC TE0752 6	1.4	3.8
Homo sapiens mRNA for KIAA446 protein; complete cds					3.8
Interferon samulated gene (2AU) Homo sapiens mRNA for KIAA46 protein; complete cds Homo sapiens mRNA for KIAA92 protein; complete cds Homo sapiens mRNA for KIAA93 for protein AD7c-NTP Haspiens]  AA714183 AA174183 J 2.0  AA27955 RC_AA47955  0.5  ESTE; Weakly similar to neuronal thread protein AD7c-NTP Haspiens]  AA27955 RC_AA427955 RC_AA427955  AA27955 RC_AA427955  AA27955 RC_AA427955  AA27955 RC_AA47955  AA27955 RC_AA47945  AA27955 RC_AA47945  AA27955 RC_AA47945  AA27956 RC_AA7745  AA27956 RC_AA77445  AA77445 RC_AA77445  AA77445 RC_AA77445  AA77445 RC_AA77445  AA77445 RC_AA77445  AA77445 RC_AA77445  AA7745 RC_AA77445  AA7745 RC_AA7745  AA7745 RC_AA7745  AA7745 RC_AA774744  AA7745 RC_AA7745  AA7745 RC_AA7745  AA7745 RC_AA7745  AA7745 RC_AA7745  AA7745 RC_AA7745  AA7745 RC_AA7745  AA7745 RC_AA77474  AA7745 RC_AA7745  AA7745 RC_AA7745  AA7745 RC_AA7745  AA7745			<b>—</b>		3.8
Homo sapiens mRNA for KNA492 protein; complete cds ESTS; Weakly similar to alternatively spliced product using exon 13A H. sapiens] ESTS; Highly similar to ZINC FINGER PROTEIN ZFP-92 [Mus musculus] ESTS; Highly similar to ZINC FINGER PROTEIN ZFP-92 [Mus ESTS; Highly similar to ZINC FINGER PROTEIN ZFP-92 [Mus musculus] ESTS; Highly similar to ZINC FINGER PROTEIN ZFP-92 [Mus ESTS; Meakly similar to neuronal thread protein AD7c-NTP [H.sapiens] ESTS; Weakly similar to neuronal thread protein AD7c-NTP [H.sapiens] ESTS; Weakly similar to neuronal thread protein AD7c-NTP [H.sapiens] ESTS; Moderately similar to neuronal thread protein AD7c-NTP [H.sapiens] ESTS; Moderately similar to neuronal thread protein AD7c-NTP [H.sapiens] ESTS; Moderately similar to neuronal thread protein AD7c-NTP [H.sapiens] ESTS; Moderately similar to neuronal thread protein AD7c-NTP [H.sapiens] ESTS; Moderately similar to neuronal thread protein AD7c-NTP [H.sapiens] ESTS; Moderately similar to neuronal thread protein AD7c-NTP [H.sapiens] ESTS; Moderately similar to neuronal thread protein AD7c-NTP [H.sapiens] ESTS; Moderately similar to neuronal thread protein AD7c-NTP [H.sapiens] ESTS; Moderately similar to neuronal thread protein AD7c-NTP [H.sapiens] ESTS; Moderately similar to neuronal thread protein AD7c-NTP [H.sapiens] ESTS; Moderately similar to neuronal thread protein AD7c-NTP [H.sapiens] ESTS; Moderately similar to neuronal thread protein AD7c-NTP [H.sapiens] ESTS; Moderately similar to neuronal thread protein AD7c-NTP [H.sapiens] ESTS; Moderately similar to neuronal thread protein AD7c-NTP [H.sapiens] ESTS; Moderately similar to neuronal thread protein AD7c-NTP [H.sapiens] ESTS; Moderately similar to neuronal thread protein AD7c-NTP [H.sapiens] ESTS; Moderately similar to neuronal thread protein AD7c-NTP [H.sapiens] ESTS; Moderately similar to neuronal thread protein AD7c-NTP [H.sapiens] ESTS; Moderately similar to neuronal thread protein AD7c-NTP [H.sapiens] ESTS; Moderately similar to neuronal thread protein AD7c-NTP [H.sapiens] EST					3.8
### Homo sapiens mRNA for KIAASP protein Complete dos procedular et al. (1985) ### RC_H99653   1.5   ### ESTs (Weakly similar to allernatively spliced product using exon 13A [H.sapiens] ### RC_T26494_f			<del></del>		3.8
### 13.4   H. sapiens   ### 13.5   H. sapiens   ### 13.5   H. sapiens   ### 13.5   ### 13.6   H. sapiens   ### 13.6   H. sapiens   ### 13.6   H. sapiens   ### 13.6   H. sapiens   ### 13.6   ### 13.6   H. sapiens   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13.6   ### 13			KC_100390_1	1.0	0.0
126494   RC_126494   RC_1264	13A [H.sapiens]	H95569	RC_H95569_i	0.5	3.7
AA774183		T26494	RC_T26494_f	1.3	3.7
cellular retunoic acid-olinding protein 1 exprint coyle membrane protein band 4.1-like 2 ESTs; Weakly similar to neuronal thread protein AD7c-NTP [H.sapiens] AA427955 RC_AA427955 RC_AA427955 RC_AA427955 RC_AA427955 RC_AA427955 RC_AA427955 RC_AA427955 RC_AA427955 RC_AA477445 RC_AA477445 RC_AA477445 RC_AA477445 RSTs; Moderately similar to neuronal thread protein AD7c-NTP [H.sapiens] RSTs AA77445 RC_AA477445 RC_AA477445 RSTs; Moderately similar to neuronal thread protein AD7c-NTP [H.sapiens] RSTs RS3769 RC_TG3769_f 1.1 RA3281PL RS376 RC_TG3769_f 1.	•	AA174183	AA174183_s	2.0	3.7
ESTS, Weakly similar to neuronal thread protein AD7c-NTP [H.sapiens] AA425378 RC_AA425378_r 0.9  WT396.31 Soares fetal liver spleen 1NFLS Homo sapiens cDNA clone IMAGE:113896 3', mRNA sequence  ESTS, Moderately similar to neuronal thread protein AD7c-NTP [H.sapiens] AA425378 RC_AA477445 RC_AA477445 RC_AA477445 RSTS; Moderately similar to neuronal thread protein AD7c-NTP [H.sapiens] AA77445 RC_AA477445 RC_AA477445 RC_AA477445 RSTS; Moderately similar to neuronal thread protein AD7c-NTP [H.sapiens] AA77485 RC_AA671089 RC_AA77445 RC_AA77445 RSTS; Moderately similar to neuronal thread protein AD7c-NTP [H.sapiens] AA77485 RC_AA77445 RC	cellular retinoic acid-binding protein 1	R53950	RC_R53950_s	1.1	3.7
RESTs; Weakly similar to neuronal thread protein AD7c-NTP   H-sapiens    A425378   RC_A425378_r   0.9		AA427955	RC_AA427955	0.5	3.7
Variable	ESTs; Weakly similar to neuronal thread protein AD7c-NTP	AA425378	RC_AA425378_r	0.9	3.7
17/525   RC_17/325   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1.5   1				4.5	2.7
ESTs; Moderately similar to neuronal thread protein AD7c-NTP [H.sapiens]	clone IMAGE:113896 3', mRNA sequence	T77525	_		3.7
		AA477445	RC_AA477445	1.4	3.7
		A A 074090	BC AA071089	1.2	3.7
Restrict					3.7
Name					3.7
Romo sapiens mRNA for Dinal protein   R2393   RC_R32393_s   1.2					3.7
Homo sapiens mRNA for KIAA878 protein; complete cds			<del>-</del>		3.7
Homo sapiens mRNA for KIAA978 protein, corriptete dus procellagen-lysine; 2-oxoglutarate 5-dioxygenase (lysine hydroxylase) 2  ESTS  ESTS  AA156335  ESTS  AA1685374  ESTS  AA1685374  ESTS  AA189375  ESTS  AA189375  ESTS  AA189375  ESTS  AA235040  ESTS  AA235040  ESTS  EST			<del>-</del>	2.8	3.7
hydroxylase) 2  ESTs  AA156335  RC_AA156335  RC_AA156335  AA156335  RC_AA156335  RC_AA156335  AA156335  RC_AA156335  AA156335  RC_AA156335  AA156335  RC_AA156335  AA156335  RC_AA156335  AA156335  RC_AA156335  AA156335  RC_AA156335  RC_AA156335  AA156335  RC_AA156335  RC_AA156335  RC_AA156335  AA156335  RC_AA156335  RC_AA15686 RC_AA15686 RC_AA15686 RC_AA15686 RC_AA15686 RC_AA15686 RC_AA156866 RC_AA16866 R		1130000	110_1100000		
ESTS  2n13d5.s1 Stratagene hNT neuron (#937233) Homo sapiens CDNA clone IMAGE:54735 3' similar to gbt.8441 CYTOCHROME C OXIDASE POLYPEPTIDE III (HUMAN); mRNA sequence fibroblast growth factor receptor 2 (bacteria-expressed kinase; keratinocyte growth factor receptor; craniofacial dysostosis 1; Crouzon syndrome; Pfeiffer syndrome; Jackson-Weiss syndrome)  ESTs; Weakly similar to eyelid [D.melanogaster]  ESTs  AA489375  AA489375  RC_AA489375_f  1.6  ESTs; Weakly similar to eyelid [D.melanogaster]  ESTs  Collagen; type III; alpha 1 (Ehlers-Danlos syndrome type IV; autosomal dominant)  ESTs; Moderately similar to !!!! ALU SUBFAMILY SC WARNING ENTRY !!!! [H.sapiens]  ESTs  R27006  RC_A235040  A7  ESTS  R27006  RC_T26471  RC_T26471  RC_T26471  RC_T26471  RC_T26471  RC_T26471  RC_T26471  RC_T26471  RC_T26471  A5  ESTS  H.sapiens mRNA for putative progesterone binding protein H60595  RC_H60595_s  1.3  TOWN SAA60450  RC_AA460450  RC_AA4		U84573			3.7
cDNA clone IMAGE:54735 3' similar to gb:L8441 CYTOCHROME C OXIDASE POLYPEPTIDE III (HUMAN); mRNA sequence fibroblast growth factor receptor; craniofacial dysostosis 1; Crouzon syndrome; Pfeiffer syndrome; Jackson-Weiss syndrome) AA489375 RC_AA489375_f 1.6 ESTs; Weakly similar to eyelid [D.melanogaster] ESTs AA235040 RC_AA235040 1.5  collagen; type III; alpha 1 (Ehlers-Danlos syndrome type IV; autosomal dominant) ESTs; Moderately similar to !!!! ALU SUBFAMILY SC WARNING ENTRY !!!! [H.sapiens] ESTs R27006 RC_R27006_f R32906 RC_R27006_f R33907 RC_T26471 RC_T26471 RC_T26471 RS_T26471	ESTs	AA156335	RC_AA156335	4.6	3.7
fibroblast growth factor receptor 2 (bacteria-expressed kinase; keratinocyte growth factor receptor; craniofacial dysostosis 1; Crouzon syndrome; Pfeiffer syndrome; Jackson-Weiss syndrome)  ESTs; Weakly similar to eyelid [D.melanogaster]  ESTs  AA489375  RC_AA489375_f  1.6  ESTs; Weakly similar to eyelid [D.melanogaster]  ESTs  AA235040  RC_AA235040  RC_AA235040  1.5  Collagen; type III; alpha 1 (Ehlers-Danlos syndrome type IV; autosomal dominant)  ESTs; Moderately similar to !!!! ALU SUBFAMILY SC WARNING ENTRY !!!! [H.sapiens]  ESTs  R27006  RC_R27006_f  RC_R27006_f  1.6  H.sapiens mRNA for putative progesterone binding protein H60595  RC_H60595_s  1.3  fibroblast growth factor receptor 2 (bacteria-expressed kinase; keratinocyte growth factor receptor; craniofacial dysostosis 1; Crouzon syndrome; Pfeiffer syndrome; Jackson-Weiss syndrome)  AA460450  RC_AA460450  RC_AA460450  RC_AA460450  1.5  ATPase; H+ transporting; lysosomal (vacuolar proton pump); alpha polypeptide; 7kD; isoform 1  ribosomal protein; large; P  AA416866  RC_AA416866_f  1.3  ESTs	cDNA clone IMAGE:54735 3' similar to gb:L8441 CYTOCHROME C OXIDASE POLYPEPTIDE III (HUMAN);,	AA085374	RC_AA085374	1.6	3.7
ESTs; Weakly similar to eyelid [D.melanogaster]  ESTs  AA235040  RC_AA235040  RC_AA235040  1.5  collagen; type III; alpha 1 (Ehlers-Danlos syndrome type IV; autosomal dominant)  ESTs; Moderately similar to !!!! ALU SUBFAMILY SC WARNING ENTRY !!!! [H.sapiens]  ESTs  R27006  RC_R27006_f  RC_R27006_f  H.sapiens mRNA for putative progesterone binding protein H60595  RC_H60595_s  RC_H60595_s  1.3  ribroblast growth factor receptor 2 (bacteria-expressed kinase; keratinocyte growth factor receptor; craniofacial dysostosis 1; Crouzon syndrome; Pfeiffer syndrome; Jackson-Weiss syndrome)  AA460450  RC_AA460450  RC_AA460450  RC_AA460450  1.5  ATPase; H+ transporting; lysosomal (vacuolar proton pump); alpha polypeptide; 7kD; isoform 1  ribosomal protein; large; P  AA416866  RC_AA416866_f  1.3  ESTs	fibroblast growth factor receptor 2 (bacteria-expressed kinase; keratinocyte growth factor receptor; craniofacial dysostosis 1;		DO 44400275 f	16	3.7
ESTs AA235040 RC_AA235040 1.5  collagen; type III; alpha 1 (Ehlers-Danlos syndrome type IV; autosomal dominant) X06700 X06700 4.7  ESTs; Moderately similar to !!!! ALU SUBFAMILY SC WARNING ENTRY !!!! [H.sapiens] T26471 RC_T26471 4.5  ESTs R27006 RC_R27006_f 1.6  H.sapiens mRNA for putative progesterone binding protein fibroblast growth factor receptor 2 (bacteria-expressed kinase; keratinocyte growth factor receptor; craniofacial dysostosis 1; Crouzon syndrome; Pfeiffer syndrome; Jackson-Weiss syndrome) AA460450 RC_AA460450 1.5  ATPase; H+ transporting; lysosomal (vacuolar proton pump); alpha polypeptide; 7kD; isoform 1 AA228122 RC_AA228122 1.1  ribosomal protein; large; P AA416866 RC_AA416866_f 1.3  ESTs R21443 R21443 1.6			<del>-</del>		3.7
collagen; type III; alpha 1 (Ehlers-Danlos syndrome type IV; autosomal dominant)  ESTs; Moderately similar to !!!! ALU SUBFAMILY SC WARNING ENTRY !!!! [H.sapiens]  ESTs  R27006  RC_R27006_f  H.sapiens mRNA for putative progesterone binding protein  fibroblast growth factor receptor 2 (bacteria-expressed kinase; keratinocyte growth factor receptor; craniofacial dysostosis 1; Crouzon syndrome; Pfeiffer syndrome; Jackson-Weiss syndrome)  AA460450  ACA4460450  RC_AA460450  RC_AA460450  1.5  ATPase; H+ transporting; lysosomal (vacuolar proton pump); alpha polypeptide; 7kD; isoform 1  ribosomal protein; large; P  AA416866  RC_AA416866_f  1.3  ESTs	· · · · · · · · · · · · · · · · · · ·		<del>-</del>		3.7
autosomal dominant)  ESTS; Moderately similar to !!!! ALU SUBFAMILY SC WARNING ENTRY !!!! [H.sapiens]  ESTS  R27006  RC_R27006_f  H.sapiens mRNA for putative progesterone binding protein H60595  RC_H60595_s  1.3  RC_H60595_s  1.3  RC_NA460450  RC_AA460450  RC_AA460450  RC_AA460450  RC_AA460450  1.5  ATPase; H+ transporting; lysosomal (vacuolar proton pump); alpha polypeptide; 7kD; isoform 1  ribosomal protein; large; P  AA416866  RC_AA416866_f		AA235040	RC_AA255040	1.0	٠١
ENTRY !!!! [H.sapiens]  ESTs  R27006  RC_R27006_f  H.sapiens mRNA for putative progesterone binding protein  fibroblast growth factor receptor 2 (bacteria-expressed kinase; keratinocyte growth factor receptor; craniofacial dysostosis 1;  Crouzon syndrome; Pfeiffer syndrome; Jackson-Weiss syndrome)  ATPase; H+ transporting; lysosomal (vacuolar proton pump); alpha polypeptide; 7kD; isoform 1  ribosomal protein; large; P  AA416866  RC_AA416866_f  RC_AA416866_f  1.3  ESTs	autosomal dominant)		X06700	4.7	3.7
ESTS R27006 RC_R27006_f 1.6  H.sapiens mRNA for putative progesterone binding protein H60595 RC_H60595_s 1.3  fibroblast growth factor receptor 2 (bacteria-expressed kinase; keratinocyte growth factor receptor; craniofacial dysostosis 1; Crouzon syndrome; Pfeiffer syndrome; Jackson-Weiss syndrome)  ATPase; H+ transporting; lysosomal (vacuolar proton pump); alpha polypeptide; 7kD; isoform 1  ribosomal protein; large; P  AA416866 RC_AA416866_f  ESTS  R27006 RC_R27006_f 1.6  RC_H60595_s 1.3  RC_AA460450 SC_AA460450  1.5  RC_AA460450 I.5  RC_AA460450 I.5  RC_AA460450 I.5  RC_AA460450 I.5  RC_AA416866_f I.3  RC_AA416866_f I.3		G T26471	RC_T26471	4.5	3.7
H.sapiens mRNA for putative progesterone binding protein  fibroblast growth factor receptor 2 (bacteria-expressed kinase; keratinocyte growth factor receptor; craniofacial dysostosis 1; Crouzon syndrome; Pfeiffer syndrome; Jackson-Weiss syndrome)  AA460450  AC_AA460450  AC_AA46		R27006	RC_R27006_f	1.6	3.7
fibroblast growth factor receptor 2 (bacteria-expressed kinase; keratinocyte growth factor receptor; craniofacial dysostosis 1; Crouzon syndrome; Pfeiffer syndrome; Jackson-Weiss syndrome)  AA460450  RC_AA460450  1.5  ATPase; H+ transporting; lysosomal (vacuolar proton pump); alpha polypeptide; 7kD; isoform 1  AA228122  RC_AA228122  1.1  ribosomal protein; large; P  AA416866  RC_AA416866_f  1.3  ESTs	i e	H60595	RC_H60595_s	1.3	3.7
ATPase; H+ transporting; lysosomal (vacuolar proton pump); alpha polypeptide; 7kD; isoform 1	fibroblast growth factor receptor 2 (bacteria-expressed kinase; keratinocyte growth factor receptor; craniofacial dysostosis 1; Crouzon syndrome; Pfeiffer syndrome; Jackson-Weiss	AA460450	RC_AA460450	1.5	3.7
alpha polypeptide; 7kD; isoform 1       AA228122       RC_AA228122       1.1         ribosomal protein; large; P       AA416866       RC_AA416866_f       1.3         ESTs       R21443       R21443       1.6	ATPase; H+ transporting; lysosomal (vacuolar proton pump);			4.4	3.7
ESTs R21443 R21443 1.6	alpha polypeptide; 7kD; isoform 1		<del></del>		ı
12175	ribosomal protein; large; P				3.7 3.7
Homo sapiens mRNA for KIAA564 protein; partial cds AA053020 RC_AA053020_I 1.0					
	Homo sapiens mRNA for KIAA564 protein; partial cds	AA053020	RC_AA053020_I	1.0	3.7

# FIGURE 8 (Cont.)

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ESTs; Highly similar to DOSAGE COMPENSATION REGULATOR [Drosophila melanogaster]	N91377	RC_N91377	2.8	3.7
ESTs	H95989	RC_H95989_s	2.0	3.7
ESTs; Weakly similar to B-cell growth factor [H.sapiens]	T88817	RC_T88817	1.0	3.7
ESTs; Weakly similar to HYPOTHETICAL 38.5 KD PROTEIN IN				-
SUI2-TDH2 INTERGENIC REGION [Saccharomyces cerevisiae]		RC_AA394126	1.8	3.6
ESTs	R38547	RC_R38547	0.9	3.6
ESTs; Weakly similar to uroporphyrinogen III synthase; UROIIIS [H.sapiens]	AA476237	RC_AA476237	1.5	3.6
ESTs	R53062	RC_R53062	0.8	3.6
ESTs; Moderately similar to alternatively spliced product using exon 13A [H.sapiens]	AA452237	RC_AA452237_i	3.6	3.6
ESTs; Weakly similar to hypothetical protein [H.sapiens]	AA040465	RC_AA040465	1.2	3.6
ESTs	AA350781	RC_AA350781	1.1	3.6
ESTs	AA121313	RC_AA121313	1.4	3.6
ESTs	AA234966	RC_AA234966	1.6	3.6
ESTs	W58461	RC_W58461	1.0	3.6
ESTs	AA252372	RC_AA252372	1.2	3.6
ESTs	W31470	RC_W31470	1.5	3.6
ESTs; Weakly similar to hypothetical protein [H.sapiens]	D80076	RC_D80076_f	1.7	3.6
ubiquitin specific protease 7 (herpes virus-associated)	R54935	RC_R54935_s	1.7	3.6
ESTs; Weakly similar to SH3BGR PROTEIN [H.sapiens]	D30930	D30930 s	1.1	3.6
Homo sapiens mRNA for low molecular mass ubiquinone-binding protein; complete cds	N77716	N77716_s	1.2	3.6
ESTs; Weakly similar to alternatively spliced product using exon 13A [H.sapiens]	R56485	RC_R56485	1.0	3.6
ESTs	AA135894	RC_AA135894	1.3	3.6
zm97f8.s1 Stratagene colon HT29 (#937221) Homo sapiens	701100007	110_01100001	1.0	0.0
cDNA clone IMAGE:545895 3', mRNA sequence	AA079487	RC_AA079487	1.5	3.6
ESTs	N22152	RC_N22152_f	1.9	3.6
ESTs	AA114893	RC_AA114893	1.2	3.6
Homo sapiens HRIHFB2115 mRNA; partial cds	AA278400	RC_AA278400_f	1.5	3.6
ESTs; Weakly similar to similar to SP:YR4_BACSU [C.elegans]	W67789	RC_W67789	1.2	3.6
ESTs; Weakly similar to cDNA EST EMBL:C1359 comes from this gene [C.elegans]	N89819	RC_N89819	1.4	3.6
ESTs; Moderately similar to (defline not available 446549)		50 11100050		
[H.sapiens]	AA488658	RC_AA488658	2.4	3.6
Human clone 121711 defective mariner transposon Hsmar2 mRNA sequence	H88535	RC_H88535_f	1.3	3.6
ESTs	AA459255	RC_AA459255	1.3	3.6
immunoglobulin gamma 3 (Gm marker)	M87789	M87789	1.2	3.6
Homo sapiens signalosome subunit 2 (SGN2) mRNA; complete				
cds	AA458919	RC_AA458919	1.2	3.6
sorting nexin 3	W49551	RC_W49551	1.2	3.6
ESTs; Highly similar to (defline not available 3915613) [H.sapiens]	W38597	W38597_s	1.1	3.6
ESTs	AA446451	RC_AA446451	1.1	3.6
hemoglobin; gamma A	H74317	RC_H74317_s	0.2	3.6
neuromedin B	X76534	X76534	2.2	3.6
ESTs; Highly similar to 26S PROTEASE REGULATORY SUBUNIT 6 [Homo sapiens]	AA441978	RC_AA441978	1.2	3.6
ESTs; Moderately similar to histone H2B [H.sapiens]	AA610040	RC_AA610040	1.1	3.6
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# FIGURE 8 (Cont.)

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Human glutamate dehydrogenase (GDH) mRNA; complete cds	T86978	RC_T86978_s	1.0	3.6
Homo sapiens mRNA for leptin receptor gene-related protein	AA393825	RC_AA393825	1.1	3.6
ESTs	R49385	RC_R49385	1.5	3.6
ESTs	W16996	W16996_s	1.1	3.6
ESTs	AA351254	RC_AA351254	0.9	3.6
ESTs; Weakly similar to neuronal tyrosine threonine phosphatase 1 [M.musculus]	T88897	RC_T88897	0.9	3.6
ESTs	N70873	RC_N70873	1.1	3.5
ESTs	AA236532	RC_AA236532_s	1.0	3.5
APOLIPOPROTEIN AI REGULATORY PROTEIN-1	AA393876	RC_AA393876_s	0.9	3.5
ESTs	AA027229	RC_AA027229	1.3	3.5
ESTs; Weakly similar to The KIAA147 gene product is related to				
adenylyl cyclase. [H.sapiens]	AA131394	RC_AA131394	1.4	3.5
ESTs	AA235505	RC_AA235505	1.4	3.5
ESTs	N21207	RC_N21207	1.6	3.5
protease; serine; 11 (IGF binding)	D87258	D87258	2.4	3.5
Homo sapiens SNC73 protein (SNC73) mRNA; complete cds	H27498	RC_H27498_f	1.1	3.5
ESTs	AA621788	RC_AA621788	1.1	3.5
Human mRNA for KIAA249 gene; complete cds	T95515	RC_T95515_s	1.6	3.5
ESTs	AA043960	RC_AA043960	1.1	3.5
ribosomal protein; large; P	W32281	RC_W32281_f	1.3	3.5
ESTs; Highly similar to POTASSIUM CHANNEL PROTEIN KV2.1 [Rattus norvegicus]	T89084	RC_T89084	1.0	3.5
ESTs	T52700	RC_T52700	0.9	3.5
ESTs	H16772	RC_H16772	1.2	3.5
ATP synthase; H+ transporting; mitochondrial F complex; subunit c (subunit 9) isoform 3	AA112059	RC_AA112059_s	1.1	3.5
ESTs; Highly similar to HYPOTHETICAL 17.2 KD PROTEIN F44E2.6 IN CHROMOSOME III [Caenorhabditis elegans]	R49920	RC_R49920	1.5	3.5
ESTs; Moderately similar to neuronal thread protein AD7c-NTP [H.sapiens]	N54909	RC_N54909_s	2.3	3.5
ESTs	AA033974	RC_AA033974	1.6	3.5
ESTs; Weakly similar to Weak similarity with Salmonella typhimurium RFBU protein [C.elegans]	AA057832	RC_AA057832	1.2	3.5
synaptophysin	R42172	RC_R42172	0.6	3.5
ESTs	W27770	W27770	0.9	3.5
EST	AA164676	RC_AA164676	1.2	3.5
ESTs; Highly similar to THROMBOXANE A2 RECEPTOR [Homo sapiens]	AA253424	RC_AA253424	1.0	3.5
ESTs; Moderately similar to !!!! ALU SUBFAMILY SQ WARNING ENTRY !!!! [H.sapiens]		- RC_N23761	1.8	3.5
GLUCOSYLCERAMIDASE PRECURSOR	T48672	RC_T48672_s	1.1	3.5
collagen; type VI; alpha 3	X52022	X52022	2.6	3.5
ESTs; Highly similar to HYPOTHETICAL 44.2 KD PROTEIN IN SCO2-MRF1 INTERGENIC REGION [Saccharomyces cerevisiae]	T35725	T35725_s	2.1	3.5
ESTs; Weakly similar to !!!! ALU SUBFAMILY SP WARNING ENTRY !!!! [H.sapiens]	T40145	T40145	1.1	3.5
Human mRNA for KiAA9 gene; complete cds	H86350	RC_H86350_s	1.3	3.5
ESTs; Moderately similar to neuronal thread protein AD7c-NTP [H.sapiens]	R81173	RC_R81173	1.3	3.5
fatty-acid-Coenzyme A ligase; long-chain 3	AA316272	AA316272	1.7	3.5
ESTs				- 1
2013	R46209	RC_R46209	1.4	3.5

# FIGURE 8 (Cont.)

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ESTs	AA397916	RC_AA397916_i	1.4	3.5
ESTs	T89379	RC_T89379	1.1	3.5
ESTs	H98714	RC_H98714_s	1.6	3.5
ESTs	N69552	RC_N69552	1.2	3.5
Human alpha satellite and satellite 3 junction DNA sequence	M21305	M21305	29.9	0.3
transcription factor AP-2 alpha (activating enhancer-binding	A A 450704	A A APOZO4 :	40.0	
protein 2 alpha)	AA458761 HG1763-HT1780	AA458761_i	13.9	0.8
Prolactin-Induced Protein		HG1763-HT1780	11.9	0.4
ESTs	AA164586	RC_AA164586_s	6.2	0.8
Human protein immuno-reactive with anti-PTH polyclonal antibodies mRNA; partial cds	AA447146	RC_AA447146_s	5.9	1.6
H4 histone family; member G	X60486	X60486	5.8	1.5
ESTs	AA424798	RC_AA424798	5.5	2.9
ESTs	AA425309	RC_AA425309	5.4	1.2
ESTs	R55185	RC_R55185	5.3	1.2
Homo sapiens mRNA for KIAA48 protein; complete cds	AA412149	RC_AA412149	5.0	2.0
ESTs; Weakly similar to neuronal thread protein AD7c-NTP				
[H.sapiens]	AA621557	RC_AA621557	5.0	1.3
desmoplakin (DPI; DPII)	W9507 <b>0</b>	RC_W95070	5.0	2.6
ESTs	R49482	RC_R49482	4.6	2.0
ESTs	AA406145	RC_AA406145_f	4.6	3.0
dual specificity phosphatase 4	U48807	U48807	4.5	0.5
ESTs; Weakly similar to !!!! ALU SUBFAMILY J WARNING ENTRY !!!! [H.sapiens]	N22107	RC_N22107	4.5	2.4
golgi SNAP receptor complex member 1	AA481414	RC_AA481414	4.4	0.9
Human protein immuno-reactive with anti-PTH polyclonal antibodies mRNA; partial cds	U28831	U28831	4.4	0.6
matrix metalloproteinase 7 (matrilysin; uterine)	L22524	L22524	4.4	0.6
Homo sapiens clone 24629 mRNA sequence	AA419386	RC_AA419386	4.3	1.3
EST	W86779	RC_W86779	4.3	0.7
ser-Thr protein kinase related to the myotonic dystrophy protein kinase	N39214	RC_N39214	4.3	0.5
ESTs	T10100	RC_T10100_f	4.3	0.9
ESTs	AA251297	RC_110100_1 RC_AA251297	4.3	2.9
mammaglobin 1	U33147	U33147	4.2	0.7
ESTs; Weakly similar to !!!! ALU CLASS B WARNING ENTRY				
[!!! [H.sapiens]	N66845	RC_N66845	4.2	0.5
Sjogren syndrome antigen A2 (6kD; ribonucleoprotein autoantigen SS-A/Ro)	AA075182	RC_AA075182	4.2	2.0
ESTs	AA411621	RC_AA411621	4.1	1.2
ESTs	AA255933	RC_AA255933	4.0	1.4
ESTs	H88496	H88496_s	4.0	1.3
Protein Kinase Ht31, Camp-Dependent	HG2167-HT2237	HG2167-HT2237	3.8	1.2
prolactin-induced protein	J03460	J03460_s	3.8	0.6
ESTs	W81552	RC_W81552	3.8	1.0
ESTs	AA398892	RC_AA398892	3.8	1.9
X-box binding protein 1	M31627	M31627	3.8	0.8
ESTs	H48032	RC_H48032	3.7	3.3
protein tyrosine phosphatase; receptor type; F	Y00815	Y00815	3.7	1.2
homolog of mouse quaking QKI (KH domain RNA binding protein)	AA280004	RC_AA280004	3.7	1.5
ESTs	F10707	RC_F10707	3.7	1.3
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# FIGURE 8 (Cont.)

** 1				
ESTs	AA255991	RC_AA255991	3.7	1.0
cadherin 3; P-cadherin (placental)	X63629	X63629	3.7	0.5
ESTs	N67149	RC_N67149	3.5	3.3
Homo sapiens CD24 signal transducer mRNA, complete cds	L33930	L33930	2.5	
and 3' region	W86835	RC_W86835	3.5	1.1
copine III	AA042990	<del>-</del>	3.5	1.9
Homo sapiens mRNA for semaphorin E; complete cds	AA042990 AA070485	RC_AA042990_s	3.5	1.0
Homo sapiens clone 23967 unknown mRNA; partial cds Homo sapiens mRNA for KIAA882 protein; partial cds	Z39762	RC_AA070485	3.4	2.6
ESTs	AA419622	RC_Z39762_s RC_AA419622	3.4 3.4	0.9 3.2
SRY (sex determining region Y)-box 4	AA458584	AA458584	3.4	0.4
Human gastrointestinal tumor-associated antigen GA733-1	A430304	747430304	3.4	0.4
protein gene, complete cds, clone 5516	J04152	J04152_rna1	3.4	0.4
ESTs; Highly similar to RING3 PROTEIN [Homo sapiens]	AA188647	RC_AA188647	3.3	2.8
ESTs; Weakly similar to Numblike [M.musculus]	AA463254	RC_AA463254_s	3.3	1.6
wee1+ (S. pombe) homolog	T16282	RC_T16282_f	3.3	1.1
SWI/SNF related; matrix associated; actin dependent regulator				ĺ
of chromatin; subfamily a; member 4	U29175	U29175	3.3	3.4
Homo sapiens mRNA for squalene epoxidase, partial cds	D78129	D78129	3.3	1.4
ESTs	AA456687	AA456687	3.3	2.2
ESTs; Weakly similar to NF-kB subunit [H.sapiens]	AA487561	RC_AA487561	3.3	1.2
hepatocyte growth factor activator inhibitor	AA157857	RC_AA157857_s	3.3	2.4
MITOCHONDRIAL 6S RIBOSOMAL PROTEIN L3	X06323	X06323	3.3	2.1
ESTs	AA496053	RC_AA496053	3.3	1.8
ESTs	AA058846	RC_AA058846	3.3	3.3
ESTs; Highly similar to N-TERMINAL ACETYLTRANSFERASE	C02582	C02582	3.3	2.2
COMPLEX ARD1 SUBUNIT HOMOLOG [Leishmania donovani] stanniocalcin 2	AA126474	RC_AA126474	3.2	3.3 0.2
STATHMIN	D51276	RC_D51276_f	3.2	3.0
collagen; type I; alpha 1	Z74615	Z74615	3.2	3.0
ESTs	AA485431	RC_AA485431_s	3.2	2.4
ESTs	H89575	H89575_s	3.2	0.6
ESTs	T17185	RC T17185	3.2	3.0
ESTs	R39044	RC_R39044	3.2	0.8
ESTs	D60411	RC_D60411_s	3.2	0.8
ESTs; Weakly similar to ubiquitous TPR motif; Y isoform	200111		J. <b>L</b>	5.0
[H.sapiens]	T91518	RC_T91518_f	3.2	2.5
3-prime-phosphoadenosine 5-prime-phosphosulfate synthase 1	AA165526	RC_AA165526	3.2	1.8
androgen receptor (dihydrotestosterone receptor; testicular				l
feminization; spinal and bulbar muscular atrophy; Kennedy disease)	M35851	M35851	3.2	10
Human mRNA for calgizzarin; complete cds	D38583	D38583	3.2	1.8 2.3
Human 26S proteasome-associated pad1 homolog (POH1)	<b>D</b> 30303	D30303	J.2	2.5
mRNA; complete cds	AA621752	RC_AA621752	3.2	2.5
heterogeneous nuclear ribonucleoprotein U (scaffold attachment factor A)	R99599	RC_R99599_s	3.2	3.4
ESTs	AA088228	RC_AA088228	3.2	1.4
ESTs	W72838	RC_W72838	3.2	1.0
ESTs	AA485212	RC_AA485212	3.1	1.4
keratin 19	Y00503	Y00503	3.1	1.1
collagen; type XV; alpha 1	L25286	L25286	3.1	3.4
ESTs; Weakly similar to predicted using Genefinder [C.elegans]	AA416886	RC_AA416886	3.1	3.1

# FIGURE 8 (Cont.)

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ESTs	AA406294	RC_AA406294	3.1	3.1
Human aipha-1 collagen type I gene, 3' end	M55998	M55998	3.1	1.7
ESTs	W02734	RC_W02734	3.1	1.4
KERATIN; TYPE II CYTOSKELETAL 7	M13955	M13955	3.1	1.1
H.sapiens mRNA for retrotransposon	AA598453	RC_AA598453_s	3.1	0.7
ESTs; Weakly similar to DREBRINS E1 AND E2 [Gallus gallus]	N69879	RC_N69879_s	3.1	1.6
ESTs	N48603	RC_N48603	3.1	1.2
ESTs; Highly similar to RSP5 PROTEIN [Saccharomyces cerevisiae]	N51488	RC N51488	3.0	1.6
ESTs; Weakly similar to 25 kDa trypsin inhibitor [H.sapiens]	N67422	RC_N67422_s	3.0	1.5
ESTs; Highly similar to 6S RIBOSOMAL PROTEIN L3A [Saccharomyces cerevisiae]	AA045365	RC_AA045365	3.0	1.7
ESTs	T32108	RC_T32108	3.0	1.1
ESTs; Weakly similar to FUN9 transcript; essential gene; similar to Schizosaccharomyces pombe unknown orf SPAC24B11.8c;		DC 44504624	2.0	2.0
GenBank Accession Number Z67757 [S.cerevisiae]	AA504631	RC_AA504631	3.0	3.0

# FIGURE 8 (Cont.)

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Accession	Title
Z29083	5T4 Oncofetal antigen
AA443962	Homo sapiens histone acetyltransferase (HBO1) mRNA, complete cds
AA055656	ESTs
D20342	transducer of erbB-2 (TOB)
AA130273	ESTs; Highly similar to DOSAGE COMPENSATION REGULATOR [Drosophila melanogaster]
U23070	Human putative transmembrane protein (nma) mRNA; complete cds
AA235448	ESTs
AA256162	ESTs
AA436880	ESTs; Moderately similar to similar to rat integral membrane glycoprotein; PIR Accession Number A467 [H.sapiens]
AA256485	ESTs

				ratio			
	UniGene ID			tumor/	90%tile	edenata	ratio tumor/
Accession	<u>L.</u>	UniGene Title		body	tumor	body	normal breast
AA126474	Hs.155223	stanniocalcin 2		72.2	722	1	1.9 39.7
U20758	Hs.313	secreted phosphoprotein 1 (osteopontin; bone sialoprotein	BCJ7	45.7 40.2	457 402	1	4
AA434329	Hs.36563	ESTs	BCY2	40.2 35.9	359	10	29.7
AA250737	Hs.72472	ESTs	DC12	34.3	411	12	5.1
X82153	Hs.83942	cathepsin K (pycnodysostosis)	BCQ3	32.2	322	1	4.7
X03635	Hs.1657	estrogen receptor 1 ESTs; Weakly similar to unknown [H.sapiens]	DCQ3	30.6	306	4	26.5
H09290	Hs.76550		BCN2	29	290	1	26.8
AA428090	Hs.26102	ESTs ESTs	00112	26.3	356	14	1
AA419547	Hs.11713	ESTs	BCO2	25.4	508	20	3
AA256485	Hs.182471	ESTs	BCX9	25.1	288	12	6.7
N67239	Hs.10760	ESTs; Weakly similar to !!!! ALU SUBFAMILY J WARNING		24.2	242	10	5.6
Z38595	Hs.125019	tumor necrosis factor (ligand) superfamily; member 10		22.8	228	9	12.4
H25836	Hs.83429	Prolactin-Induced Protein		22.7	760	34	1.4
HG1763-HT1780	Hs.3838	serum-inducible kinase		22.6	226	10	0.9
C01714	нs.3636 Hs.182225	RNA binding motif protein 3		22.1	221	9	17.8
U28686		ESTs		21.2	212	6	17.4
AA411621	Hs.8895	ESTs	BCX6	20.9	209	1	19.5
N46252	Hs.29724	fetal Alzheimer antigen	DOMO	20.6	206	4	19.1
U05237	Hs.99872	dual specificity phosphatase 4		20.2	202	5	
U48807	Hs.2359	ESTs; Weakly similar to !!!! ALU SUBFAMILY SQ WARNI		18.7	187	1	
AA070801	Hs.51615	Human protein immuno-reactive with anti-PTH polyclonal		18.6	186	10	
U28831	Hs.44566			17.5	175	2	
AA292066	Hs.240802	ESTs; Weakly similar to Br140 [H.sapiens] secreted frizzled-related protein 4	BCX2	17.4	409	24	
AA291725	Hs.105700	•	BCY5	17.4	174	9	
N26722	Hs.42645	ESTs	50.0	16.6	166	8	
AA256323	Hs.25264	ESTS		16.2	162	1	
AA065217	Hs.169674	ESTs		16	255	16	
AA446650	Hs.27860	ESTs osteoblast specific factor 2 (fasciclin I-like)	BCA4	15.7	1030	66	
D13666	Hs.136348	ESTs	BCX8	15.6	156	7	
AA621169	Hs.8687	"Human neuropeptide Y receptor Y1 (NPYY1) mRNA, exc		15.3	153	1	
L07615	11- 240400	ESTs	•	15.2	152	1	
AA456598	Hs.240190	ESTs		14.9	149	1	
AA007234	Hs.30098 Hs.14838	ESTs	всх4		219	15	
F01831 N66818	нs. 14030 Hs.42179	ESTs	BCY6		145	1	
		"Protein Kinase Ht31, Camp-Dependent"		14.4	144	g	
HG2167-HT2237 Z39821	Hs.107295	ESTs		14.3	143		
H05509	Hs.24639	ESTs		14.2	142		
T90621	Hs.109052	chromosome 14 open reading frame 2		14.2	142		9.4
AA171913	Hs.5338	carbonic anhydrase XII		14.2	390	28	22.5
AA149007	Hs.243954	ESTs		13.7	137		8.9
N22222	110.210001	"yw34b06.s1 Morton Fetal Cochlea Homo sapiens cDNA		13.5	135		5.1
AA480975	Hs.44829	ESTs; Moderately similar to !!!! ALU SUBFAMILY SQ WA		13.3	133		
D62633	Hs.8236	ESTs		13.3	445	34	6
D12485	Hs.11951	phosphodiesterase I/nucleotide pyrophosphatase 1 (hom	o BCA2		244	19	9.9
AA490262	Hs.15485	ESTs; Moderately similar to APXL gene product [H.sapie			331	25	12.4
W93640	Hs.4779	ESTs		13.1	131	1	5.1
D49396	Hs.75454	Human mRNA for Apo1_Human (MER5(Aop1-Mouse)-lik	е	12.8	128		11.7
H94892	Hs.6906	v-ral simian leukemia viral oncogene homolog A (ras rela		12.8	141		
AA458761	Hs.18387	ESTs		12.7	311		
AA436158	Hs.190013	ESTs		12.6	126		7.5
AA444369	Hs.177537	ESTs		12.6	126		
X14787	Hs.87409	thrombospondin 1		12.6	126		10.8
T40327	Hs.80680	ESTs		12.5	156		3 2.9
Z48633	Hs.6940	H.sapiens mRNA for retrotransposon		12.4	124		
AA227219	Hs.110826	Homo sapiens CAGF9 mRNA; partial cds		12.3	123	. 1	11.3
T97307	Hs.161720	ESTs; Moderately similar to !!!! ALU SUBFAMILY J WAR	N.	12.3	129	11	11.7

M86849		"Homo sapiens connexin 26 (GJB2) mRNA, complete cds CBC	C2	12	120	8	9
AA417152	Hs.5101	ESTs; Highly similar to protein regulating cytokinesis 1 [H. CQ/	A4	11.8	201	17	19.1
D31352	Hs.31433	ESTs; Weakly similar to !!!! ALU SUBFAMILY SQ WARNI		11.7	117	1	10.1
AA251089	Hs.94576	ESTs; Weakly similar to phosducin; retinal [H.sapiens]		11.5	115	1	6.9
AA224180	Hs.187579	ESTs; Weakly similar to !!!! ALU SUBFAMILY J WARNING		11.5	115	1	10
F11019	Hs.12696	ESTs		11.4	114	1	10
L19872	Hs.170087	aryl hydrocarbon receptor		11.3	113	8	3.9
Al471525	Hs.97496	YY1 transcription factor		11.3	124	11	9.7
AA487557	Hs.10706	ESTs; Weakly similar to (defline not available 3882221) [H		11.3	113	8	2.5
M24594	Hs.20315	interferon-induced protein 56		11.2	112	8	5.9
AA279112	Hs.88594	ESTs		11.2	112	1	10.3
AA490969	Hs.168147	ESTs CQ	8A9	11	187	17	10.4
X17059	Hs.155956	N-acetyltransferase 1 (arylamine N-acetyltransferase)		10.8	706	66	9.2
W85765	Hs.30504	ESTs		10.7	123	12	7
AA405569	Hs.418	fibroblast activation protein; alpha CZ	Ά9	10.7	433	41	7.2
N31952	Hs.167531	ESTs; Weakly similar to (defline not available 3875448) [C		10.5	105	4	7.1
H93575	Hs.227146	ESTs		10.5	105	1	9.9
F03969	Hs.16940	ESTs; Weakly similar to tumorous imaginal discs protein T		10.5	105	1	9
N22157	Hs.226573	Homo sapiens IkB kinase-b (IKK-beta) mRNA; complete c		10.5	121	12	1.6
F13673	Hs.99769		N4	10.4	880	85	5.3
AA131692	Hs.26204	ESTs		10.3	103	1	3.9
AA411745	Hs.239681	ESTs; Weakly similar to KIAA0554 protein [H.sapiens]		10.3	103	1	9.3
D86957	Hs.80712	Human mRNA for KIAA0202 gene; partial cds		10.2	102	1	4.8
AA406542	Hs.71520	ESTs		10.2	506	50	2.8
U65932	Hs.81071		3C3	10.2	628	62	17.2
AA236813	Hs.72324	ESTs; Highly similar to unknown [H.sapiens]		10.1	111	11	10.2
M90516	Hs.1674	glutamine-fructose-6-phosphate transaminase 1		10	100	1	7.6
	Hs.33287		CQ1	9.9	483	49	1.8
AA425309	Hs.100686	ESTs; Moderately similar to secreted cement gland protei BC	CX3	9.9	351	36	13.9
AA487468	Hs.5085	dolichyl-phosphate mannosyltransferase polypeptide 1; ca		9.8	123	13	5
AF007875	Hs.45231	ESTs		9.8	98	1	8.8
AA398913		"Basic Transcription Factor, 44 Kda Subunit"		9.7	97	10	7.2
HG3748-HT401		sine oculis homeobox (Drosophila) homolog 1		9.7	97	1	9.3
X91868	Hs.54416 Hs.154554	ESTs; Weakly similar to ANKYRIN; BRAIN VARIANT 1 [H		9.7	102	11	6
AA599267		RAS p21 protein activator (GTPase activating protein) 1		9.6	96	1	8.5
M23379	Hs.758	ESTs BC	CY9	9.6	124	13	9
T25867	Hs.7549 Hs.75760	sterol carrier protein 2		9.5	95	4	8.8
U11313	Hs.110488	ESTs		9.5	95	1	8.5
R63542	Hs.620	bullous pemphigoid antigen 1 (230/240kD)		9.4	94	1	0.3
M69225		ESTs		9.4	94	8	7.3
AA250775	Hs.87747	ESTs		9.4	94	3	5.3
AI039722	Hs.171205	sialyltransferase		9.3	93	4	3
U14550	Hs.107573 Hs.159627	Death associated protein 3		9.3	93	5	8
U18321		uracil-DNA glycosylase		9.3	93	8	8.2
X89398	Hs.78853	chromosome-associated polypeptide C		9.3	93	1	8.4
AA283006	Hs.50758	MAD (mothers against decapentaplegic; Drosophila) hom		9.3	93	1	7.8
U44378	Hs.75862 Hs.21941		AD1	9.3	436	47	5.8
AA187490		ESTs		9.2	234	26	16.8
AA487202	Hs.17962	ESTs		9.1	91	6	8.3
T33637	Hs.6841	to the state of the state o		9.1	91	1	7.6
AA235112	Hs.106227	Human beige-like protein (BGL) mRNA; partial cds		9	144	16	13
M83822	Hs.62354 He 181104			9	117	13	10.6
AA256680	Hs.181104		CX5	9	90	1	5.5
AA028028	Hs.61460 Hs.239489	2010		8.9	89	5	8
M77142				8.8	110	13	10.5
AA858097	Hs.173594 Hs.73625	RAB6 interacting; kinesin-like (rabkinesin6)		8.8	199	23	16.1
AA179845	Hs.44276	ESTs; Moderately similar to HOMEOBOX PROTEIN HOX		8.7	247	29	5.7
AA112396		"Human chondroitin sulfate proteoglycan versican V splice		8.6	568	66	22.4
U16306	Hs.81800	"Epican, Alt. Splice 1"		8.5	85	1	3.2
HG2981-HT31				8.5	127	15	1.6
AA280036	Hs.145374	LO15, Medicy chilinal to 115 mais [o.ologuno]					

D83004	Hs.75355	ubiquitin-conjugating enzyme E2N (homologous to yeast U	8.5	85	1	7.2
AA609200	Hs.162686	ESTs BCY4	8.5	85	1	4.3
U33147	Hs.46452	mammaglobin 1	8.5	2058	243	1.4
N30856	Hs.30246	ESTs	8.4	84	1	5.3
AA446887	Hs.42911	ESTs	8.4	101	12	8.7
	Hs.54629	ESTs	8.4	84	10	0.8
N90526		transcription factor COUP 2 (chicken ovalbumin upstream	8.4	169	20	4.6
AA393876	Hs.1255	ESTs	8.3	83	3	1.8
AA257971	Hs.21214	ESTs	8.3	83	8	1.9
D60799	Hs.169391	v-kit Hardy-Zuckerman 4 feline sarcoma viral oncogene h	8.3	87	11	0.4
AA143045	Hs.81665	· · · · · · · · · · · · · · · · · · ·	8.3	145	18	3.7
AA047896	Hs.49169	ESTs	8.2	82	1	6.8
U59863	Hs.146847	TRAF family member-associated NFKB activator	8.2	124	15	11.5
H95094	Hs.75187	KIAA0016 gene product	8.2	82	1	7.4
H13108	Hs.107968	ESTs		114	14	9.9
AA236324	Hs.92381	ESTs; Weakly similar to !!!! ALU CLASS A WARNING EN CVA1	8.2		48	0.9
L22524	Hs.2256	matrix metalloproteinase 7 (matrilysin; uterine)	8.2	396		
T16387	Hs.65328	ESTs	8.2	82	1	6.4
W42451	Hs.92260	high-mobility group protein 2-like 1	8.1	94	12	6.5
Z38501	Hs.8768	ESTs; Weakly similar to neuronal thread protein AD7c-NT	8.1	81	9	5.5
W04517	Hs.18442	ESTs	8.1	81	3	2.8
D31161	Hs.68613	ESTs	8.1	81	1	4.6
AA452000	Hs.94030	ESTs	8.1	101	13	7.9
R40057	Hs.112360	prominin (mouse)-like 1 CZA8	8.1	328	41	1.7
AA451992	Hs.247127	ESTs; Weakly similar to similar to Schizosaccharomyces p	8	84	11	6.3
AA227428	Hs.9728	ESTs; Weakly similar to KIAA0512 protein [H.sapiens]	8	80	6	7.3
AA620599	Hs.24766	ESTs	8	100	13	2.9
N32919	Hs.27931	ESTs	7.9	79	1	6.2
AA398155	Hs.97600	ESTs	7.9	79	1	2.7
R79723	Hs.69997	H.sapiens mRNA for translin associated zinc finger protein	7.9	234	30	18.9
Z29083	Hs.82128	5T4 oncofetal trophoblast glycoprotein BCA7	7.9	79	2	6.9
Al267886	Hs.148027	polymerase (RNA) II (DNA directed) polypeptide B (140kD	7.8	137	18	11.9
M28213	Hs.78305	RAB2; member RAS oncogene family	7.8	78	1	5.6
R56678	Hs.88959	ESTs; Weakly similar to !!!! ALU SUBFAMILY SP WARNIN	7.7	. 77	8	6.9
AA031357	Hs.31803	ESTs	7.7	77	1	5.1
N66857	Hs.14808	ESTs; Weakly similar to !!!! ALU CLASS C WARNING EN	7.7	77	1	5
Z39436	Hs.102720	ESTs	7.7	81	11	7.6
T90037	Hs.16686	ESTs	7.6	76	1	4.2
	Hs.62349	ESTs	7.6	92	12	1.4
AA167268		KIAA0480 gene product	7.6	76	1	5
R34531	Hs.243068	ESTs	7.6	144	19	13.9
AA416997	Hs.59622	ESTs	7.5	112	15	2.5
AA211400	Hs.193172		7.5	75	1	6.5
D60237	Hs.14368	SH3-binding domain glutamic acid-rich protein like  FSTs PAA9	7.5	136	18	3.4
W37145	Hs.30029	2010	7.4	74	1	6
AA054228	Hs.23165	ESTs	7.4	74	3	1.7
AA455875	Hs.227602	Homo sapiens mRNA for KIAA0727 protein; partial cds	7.4	74	8	6
AA043562	Hs.62637	ESTs	7.4	103	14	6.5
D62657	Hs.35086	ubiquitin-specific protease 1	7.4	74	5	2.4
AA044842	Hs.95260	ESTs	7.4	137	19	1.8
AA159181	Hs.184013		7.3	73	1	5.3
M99701	Hs.95243	transcription elongation factor A (SII)-like 1			1	5.2
Z46629	Hs.2316	SRY (sex-determining region Y)-box 9 (campomelic dyspl	7.3	73 73	1	3.8
AA165333	Hs.24808	ESTs	7.3	73		
N90719	Hs.94445	ESTs	7.3	73	3 15	5.4 5
L38608	Hs.10247	activated leucocyte cell adhesion molecule	7.3	106	15	5
R87834	Hs.3688	acid-inducible phosphoprotein	7.3	73	1	1.2
AA042990	Hs.171921	sema domain; immunoglobulin domain (Ig); short basic do	7.3	271	37	2.3
N64378	Hs.13149	ESTs; Weakly similar to ARI protein [D.melanogaster]	7.2	72	10	6.1
AA478446	Hs.69559	ESTs; Weakly similar to Bat2 [H.sapiens]	7.2	72	1	5.7
U83908	Hs.247134		7.2	72	1	5.8
W60913	Hs.30738	ESTs	7.2	72	4	5.7

AA393164	Hs.97644	mammaglobin 2	7.	2 498	69	9.3
F10577	Hs.70312	ESTs; Moderately similar to neuronal thread protein AD7c	7.		9	6.9
AA211941	Hs.109643	polyadenylate binding protein-interacting protein 1	7.		1	6.2
X57985	Hs.2178	H2B histone family; member Q	7.		14	7.5
M57230	Hs.82065	interleukin 6 signal transducer (gp130; oncostatin M recep	7.		4	6.4
D61676	Hs.21851	Homo sapiens mRNA; cDNA DKFZp586J2118 (from clone E				3.6
H18027	Hs.184697		3CY7 7.		21	14.5
AA199828	Hs.188662	· ·	3CY8 7.		1	6.5
AA032147	Hs.23296	ESTs	7		1	6.5
AA436244	Hs.17240	ESTs	-		3	1.3
AA400080	Hs.97774	EST	;		1	0.9
U25435	Hs.57419	transcriptional repressor	-			5.4
M74524	Hs.80612	ubiquitin-conjugating enzyme E2A (RAD6 homolog)	;		14	7.5
W47183	Hs.153468	ESTs; Weakly similar to !!!! ALU SUBFAMILY SB2 WARN	-		6	1.5 6
M60752	Hs.121017	H2A histone family; member A	6.			
J03460	Hs.99949	prolactin-induced protein				8.4
AA292701	Hs.5364	ESTs	6.			1.3
AA219699		ESTs	6.		1	4.4
H64938	Hs.184245	ESTs	6.		5	6.2
	Hs.38331		6.		10	2.4
Z38839	Hs.125019	ESTs; Weakly similar to !!!! ALU SUBFAMILY J WARNING	6.		47	2.1
W86779	Hs.241582	EST	6.			2.6
AA261852	Hs.192905	ESTs	6.		1	0.2
Al283493	Hs.75722	ribophorin II	6.		33	2.8
H17861	Hs.17767	ESTs	6.		19	12.1
J04088	Hs.156346	topoisomerase (DNA) II alpha (170kD)	6.	<b>8</b> 68	1	5.6
AA608955	Hs.109653	ESTs	6.	<b>8</b> 68	10	6.1
U39840	Hs.105440	hepatocyte nuclear factor 3; alpha	6.	<b>7</b> 67	9	6.3
AA425367	Hs.32094	ESTs	6.	7 94	14	8
H48502	Hs.28212	ESTs	6.	<b>7</b> 78	12	3
Z38763	Hs.15740	ESTs	6.	<b>7</b> 67	1	6.3
AA598803	Hs.111496	ESTs	6.	<b>7</b> 67	2	2.1
Al287461	Hs.164950	ESTs	6.	7 67	1	6
N45219	Hs.48320	ESTs	6.	7 155	23	1.4
AA195260	Hs.204151	ESTs; Moderately similar to !!!! ALU SUBFAMILY SX WAR	6.	7 67	1	5.7
F09012	Hs.181326	ESTs	6.	7 67	6	1.9
L11066	Hs.3069	heat shock 70kD protein 9B (mortalin-2)	6.	7 93	14	8.4
AA453783	Hs.76550	ESTs; Weakly similar to unknown [H.sapiens]	6.	7 304	46	7.8
T25508	Hs.81057	ESTs	6.	<b>7</b> 67	9	5.7
L40391	Hs.6445	Homo sapiens (clone s153) mRNA fragment	6.	6 135	21	13.1
AA147719	Hs.159441	ESTs	6.	<b>6</b> 66	1	5.4
AA126433	Hs.173242	sorting nexin 4	6.		11	6.3
M21305	Hs.247946	Human alpha satellite and satellite 3 junction DNA sequen	6.		135	0.8
AA041551	Hs.48644	ESTs	6.		2	6
R42036	Hs.6763	ESTs	6.	<b>5</b> 65	10	1.5
T40530	Hs.231577	ESTs	6.	<b>5</b> 65	6	4.8
N29888	Hs.169539	ESTs	6.		4	5.3
AA490862	Hs.55901	ESTs; Moderately similar to !!!! ALU SUBFAMILY SQ WA	6.		1	5.6
R99599	Hs.103804	heterogeneous nuclear ribonucleoprotein U (scaffold attac		5 162	25	14.7
M63256	Hs.75124	cerebellar degeneration-related protein (62kD)	6.	-	2	4.9
U37519	Hs.87539	aldehyde dehydrogenase 8	6.		67	2.3
AB002367	Hs.21355	doublecortin and CaM kinase-like 1	6.		8	3
AA284755	Hs.214742	CDW52 antigen (CAMPATH-1 antigen)	6.		8	6
AA243012	Hs.75928	ESTs	6.		11	5
Al356250	Hs.4779	ESTs	6.		12	6.6
X06700	Hs.119571	collagen; type III; alpha 1 (Ehlers-Danios syndrome type I	6.		175	5
U51166	Hs.173824	thymine-DNA glycosylase	6.		16	4.4
W23625	Hs.8739	ESTs	6.		1	5.1
M15796	Hs.78996	proliferating cell nuclear antigen	6.		39	22.4
HG4390-HT4660		Ribosomal Protein L18a Homolog	6.		4	5.7
X92098	Hs.75914	H.sapiens mRNA for transmembrane protein rnp24	6.		16	9.1
		, protein nipa T	0.	- 30	10	9.1

1107744		III.					
N67711	Hs.151046	Homo sapiens clone 23859 mRNA sequence		6.3	63	1	5.8
W37999	Hs.24336	ESTs		6.3	63	6	5
AA149894	Hs.20815	erythroblast macrophage protein		6.3	165	26	3.2
H10933	Hs.10067	ESTs	BCA1	6.3	693	110	7.2
AA609723	Hs.30652	ESTs	PAA3	6.3	63	1	5.4
AA122386	Hs.82985	collagen; type V; alpha 2		6.3	1075	171	3.8
AA243052	Hs.172643	Homo sapiens mRNA; cDNA DKFZp564J1616 (from clone		6.2	62	6	5.6
R39995	Hs.25925	Homo sapiens clone 23860 mRNA sequence		6.2	62	2	5.9
AA430124	Hs.234607	ESTs		6.2	62	1	5.4
T68510	Hs.76704	ESTs		6.2	600	97	4.1
R79750	Hs.83623	constitutive androstane receptor-beta; orphan nuclear hor		6.1	493	81	0.7
U35835	Hs.155637	protein kinase; DNA-activated; catalytic polypeptide		6.1	61	1	5.7
AA046405	Hs.5306	ESTs; Weakly similar to KIAA0597 protein [H.sapiens]		6.1	61	2	5.9
AA358268	Hs.95464	ESTs; Moderately similar to transcription factor RTEF-1 [H		6.1	61	1	5.1
D31058	Hs.24375	ESTs	CXA3	6.1	343	56	16.4
U70322	Hs.168075	karyopherin (importin) beta 2		6.1	126	21	2.4
R46025	Hs.7413	ESTs		6.1	185	31	6.6
W68845	Hs.24095	ESTs		6.1	110	18	10.2
AA176690	Hs.4084	ESTs		6	60	6	4.6
N67390	Hs.43228	ESTs		6	60	5	3.7
L09717	Hs.8262	lysosomal-associated membrane protein 2		6	60	5	5.9
F03819		ESTs		6			
	Hs.173094				202	34	3.7
D38491	Hs.247463	Human mRNA for KIAA0117 gene; partial cds		5.9	59	1	2.6
F02582	Hs.14474	ESTs		5.9	59	10	4.2
AA347193	Hs.62180	ESTs	_	5.9	59	1	4.2
AA504642	Hs.28436	ESTs; Weakly similar to coded for by C. elegans cDNA CE	=	5.9	59	1	4.4
AA476594	Hs.9817	arg/Abl-interacting protein ArgBP2		5.9	186	32	3.7
Z81326	Hs.78589	protease inhibitor 12 (neuroserpin)		5.9	59	1	3.3
F10707	Hs.181104	ESTs		5.9	208	36	1.8
X07696	Hs.80342	keratin 15		5.8	753	131	0.4
X53793	Hs.117950	multifunctional polypeptide similar to SAICAR synthetase		5.8	218	38	13
AB000221	Hs.16530	small inducible cytokine subfamily A (Cys-Cys); member 1		5.8	58	1	3.2
AA458904	Hs.26267	ESTs; Weakly similar to torsinA [H.sapiens]		5.8	58	5	3.1
W63793	Hs.75744	S-adenosylmethionine decarboxylase 1		5.8	151	26	11.4
AA262491	Hs.186572	ESTs		5.8	58	1	5
AA429038	Hs.40541	ESTs		5.8	58	1	4.4
AA608531	Hs.170313	ESTs		5.8	58	1	4.9
L19161	Hs.211539	eukaryotic translation initiation factor 2; subunit 3 (gamma		5.8	171	30	2.9
R27296	Hs.23240	ESTs		5.8	115	20	2.5
AA610086	Hs.32990	ESTs		5.8	91	16	1.4
D87685	Hs.78893	Human mRNA for KIAA0244 gene; partial cds		5.8	58	1	4.9
AA262943	Hs.23552	ESTs	BCH5	5.8	336	58	2
D60302	Hs.108977	ESTs	BCY1	5.8	321	55	17
AA194882	Hs.19522	ESTs		5.7	57	8	5.3
AA287097	Hs.244443	transcription factor 4		5.7	57	8	4.1
AA463745	Hs.29403	ESTs; Weakly similar to PROBABLE ATP-DEPENDENT R	<u>:</u>	5.7	57	10	4.8
AA490814	Hs.24170	ESTs	•	5.7	94	17	7.3
AA084677	Hs.54558	ESTs; Weakly similar to !!!! ALU CLASS C WARNING EN		5.7	57	1	4.9
Z39301	Hs.7859	ESTs		5.7	57	1	4.9
AA421562	Hs.91011	anterior gradient 2 (Xenopus laevis; secreted cement glan		5.7	368	65	28.5
AA044095	Hs.3402	ESTs		5.7	57	1	4.5
AA092376	Hs.90606	15 kDa selenoprotein					
R51309	Hs.70823	KIAA1077 protein	BCN5	5.7	57 567	100	5 67
X16396	Hs.154672	methylene tetrahydrofolate dehydrogenase (NAD+ depen	DCNO	5.7	567 251	100	6.7
AA024835				5.7	251	44	6.6
D87448	Hs.47584 Hs.91417	potassium voltage-gated channel; delayed-rectifier; subfa		5.7	85 76	15	7.8
U90914		Homo sapiens mRNA for DNA topoisomerase II binding p		5.6	76 50	14	2
	Hs.5057	carboxypeptidase D		5.6	56	1	5.3
X72841 AA281591	Hs.31314	H.sapiens IEF 7442 mRNA		5.6	191	34	3.5
AA464428	Hs.16193	Homo sapiens mRNA; cDNA DKFZp586B211 (from clone		5.6	101	18	1.6
7V4U4420	Hs.119394	ESTs		5.6	108	20	1.8

AA280617	Hs.100861	ESTs; Weakly similar to p60 katanin [H.sapiens]		5.6	73	13	6.1
X52003	Hs.1406	trefoil factor 1 (breast cancer; estrogen-inducible sequenc BC		5.6	1346	239	5.4
AA121266	Hs.34641		AA8	5.6	95	17	9.1
AA521472	Hs.73435	ESTs		5.6	106	19	9
D87469	Hs.57652	EGF-like-domain; multiple 2		5.5	145	27	2.2
AA452411	Hs.29679	ESTs		5.5	147	27	4.4
AA504631	Hs.26813	ESTs; Weakly similar to (defline not available 4689108) [H		5.5	130	24	12.5
AA621557	Hs.58633	ESTs; Moderately similar to !!!! ALU SUBFAMILY SQ WA		5.5	315	58	3.1
AI051602	Hs.4112	acetyl-Coenzyme A acetyltransferase 2 (acetoacetyl Coen		5.5	55	10	4.2
AA418069	Hs.241391	natural killer-tumor recognition sequence		5.5	63	12	1
T26989	Hs.121576	aspartate beta-hydroxylase		5.5	79	15	4.4
AA143019	Hs.182667	ESTs; Highly similar to surface 4 integral membrane prote		5.5	90	17	2.9
W90146	Hs.35962	ESTs		5.5	168	31	4.4
N37065	Hs.44856	ESTs; Moderately similar to !!!! ALU SUBFAMILY SQ WA		5.5	323	59	10.5
T23983	Hs.7365	ESTs; Moderately similar to !!!! ALU SUBFAMILY SQ WA		5.5	186	34	16.5
AA130273	Hs.7584	ESTs; Weakly similar to (defline not available 4240269) [H Bo	CF3	5.5	55	1	5.2
AA262942	Hs.79741	ESTs		5.5	343	62	2.5
M86546	Hs.155691	pre-B-cell leukemia transcription factor 1		5.4	180	34	15.9
U15932	Hs.2128	dual specificity phosphatase 5		5.4	137	26	2.5
AA338760	Hs.15159	ESTs		5.4	54	1	4.4
AA460350	Hs.22370	ESTs		5.4	75	14	0.8
AA133250	Hs.62180	ESTs		5.4	54	1	4
N22414		"yw39a07.s1 Weizmann Olfactory Epithelium Homo sapie		5.4	54	1	3.7
N63823	Hs.220470	ESTs; Moderately similar to !!!! ALU SUBFAMILY SQ WA		5.4	54	1	4.3
W60473	Hs.57787	ESTs		5.4	54	1	4.1
AA287115	Hs.99697	ESTs		5.4	54	10	2.5
AA313639	Hs.185783	ESTs		5.4	73	14	6.8
M14219	Hs.76152	decorin		5.4	144	27	13.3
N79 <b>749</b>	Hs.87627	ESTs		5.4	81	15	2.6
H895 <b>75</b>	Hs.93468	ESTs		5.4	259	48	1.4
D59894	Hs.34782	ESTs BO	CJ1	5.4	483	90	4
AA485223	Hs.34892	ESTs		5.4	192	36	4.4
U43189	Hs.82143	Human Ets transcription factors NERF-1a and NERF-1b (		5.3	53	1	4.8
AA347973	Hs.221132	ESTs		5.3	67	13	5.3
AA459657	Hs.12311	Homo sapiens clone 23570 mRNA sequence		5.3	166	32	14.9
AA130596	Hs.71331	ESTs; Weakly similar to potent heat-stable protein phosph		5.3	53	1	2.8
R45175	Hs.117183	ESTs		5.3	53	6	2.3
Z39549	Hs.153746	ESTs		5.3	53	5	0.9
AA292655	Hs.96557	ESTs		5.3	58	11	3.3
F02641	Hs.12342	Homo sapiens clone 24538 mRNA sequence		5.3	53	9	3.6
AA610070	Hs.151469	ESTs; Highly similar to CASK [H.sapiens]		5.3	53	9	3.2
W52493	Hs.13531	ESTs		5.3	53	1	4.1
H46617	Hs.172241	"yp19h1.r1 Soares breast 3NbHBst Homo sapiens cDNA C\	VA3	5.3	144	27	13.1
AA449887		ESTs		5.2	52	1	4.3
AA101416	Hs.107149	ESTs		5.2	52	3	2.3
AA406546	Hs.71968	ESTs		5.2	405	78	10.1
AA465701	Hs.108646	ESTs		5.2	52	4	3.9
W80702	Hs.58461	ESTs		5.2	52	6	1.8
AA837495	Hs.69851	ESTs; Weakly similar to Wiskott-Aldrich syndrome protein		5.2	81	16	1.1
AA090695	Hs.181385	ESTs		5.2	75	15	6.4
AA132007	Hs.167420	ESTs		5.2	251	48	21
Y00503	Hs.182265	keratin 19		5.2	1320	256	3.2
AA418230	Hs.8172	ESTs; Weakly similar to alternatively spliced product using		5.2	52	1	4.9
M22995	Hs.865	RAP1A; member of RAS oncogene family		5.2	52	1	3
U89326	Hs.87223	bone morphogenetic protein receptor; type IB		5.2	52	5	3.5
X63629	Hs.2877	cadherin 3; P-cadherin (placental)		5.2	331	64	1.5
AA122147	Hs.64691	Homo sapiens mRNA for KIAA0483 protein; partial cds		5.2	117	23	5
D14878	Hs.82043	D123 gene product		5.1	106	21	9.2
AA236559	Hs.8768	ESTs; Weakly similar to neuronal thread protein AD7c-NT		5.1	181	36	15.8
AA598710	Hs.23740	ESTs		5.1	298	59	4.4

4.4050000							
AA252863	Hs.87729	ESTs		5.1	51	1	4.2
AA456099	Hs.176376	ESTs		5.1	51	1	2
AA479362	Hs.47144	ESTs		5.1	198	39	17.9
AA398302	Hs.127437	ESTS		5.1	51	1	2.4
W68502	Hs.180201	ESTs; Weakly similar to !!!! ALU CLASS C WARNING EN		5.1	58	12	5.7
R51273	Hs.79029	ESTs		5.1	51	9	3.8
N46086	Hs.92308	ESTs		5.1	150	30	7.2
N33236	Hs.28555		3CO1	5.1	51	1	3.9
M22898	Hs.1846	tumor protein p53 (Li-Fraumeni syndrome)		5.1	97	19	9.3
W44735	Hs.9286	ESTs		5.1	51	5	4.5
Z39053	Hs.27263	ESTs		5.1	113	22	6.1
U67319	Hs.9216	caspase 7; apoptosis-related cysteine protease		5	66	13	5.3
AA004415	Hs.106106	ESTs		5	468	94	4.7
F10770	Hs.180378	Homo sapiens clone 669 unknown mRNA; complete sequ		5	50	5	4.1
N26101	Hs.7838	ESTs		5	50	1	4.3
N36421	Hs.107854	ESTs; Moderately similar to !!!! ALU SUBFAMILY SQ WA		5	50	1	4.7
N57773	Hs.93560	ESTs; Weakly similar to Similar to Rat trg gene product [C		5	50	2	3.1
N93839	Hs.39288	ESTs; Weakly similar to !!!! ALU SUBFAMILY SB WARNIN		5	50	5	4.7
AA608679	Hs.108327	damage-specific DNA binding protein 1 (127kD)		5	121	25	5.9
Al382972	Hs.82128	5T4 oncofetal trophoblast glycoprotein		5	285	58	1.2
W78968	Hs.241880	H3 histone; family 3A		5	264	53	3.4
AA129465	Hs.106843	ESTs; Weakly similar to hypothetical protein [H.sapiens]		5	50	1	3.3
H88033	Hs.109727	Homo sapiens mRNA for KIAA0733 protein; partial cds		5	64	13	6.3
U84573	Hs.41270	procollagen-lysine; 2-oxoglutarate 5-dioxygenase (lysine h		5	225	45	9.1
N59764	Hs.5398	GUANINE-MONOPHOSPHATE SYNTHETASE		5	50	1	4.1
H06195	Hs.7194	ESTs		5	110	22	9.7
U47414	Hs.79069	cyclin G2		5	50	1	3.2
HG3510-HT3704		V-Erba Related Ear-3 Protein		5	82	17	0.9
U24576		LIM domain only 4		4.9	49	1	3.6
X65724	Hs.2839	Norrie disease (pseudoglioma)		4.9	49	5	3.8
X98263	Hs.152720	M-phase phosphoprotein 6		4.9	153	31	2.4
AA315807	Hs.106227	ESTs; Weakly similar to (defline not available 4200325) [H		4.9	49	1	4.2
AA348014	Hs.23412	ESTs		4.9	49	1	4.5
AA446949	Hs.6236	ESTs		4.9	337	70	2.7
AA456981	Hs.35349	ESTs		4.9	49	1	4.1
AA193592	Hs.42300	ESTs; Moderately similar to !!!! ALU SUBFAMILY SQ WA		4.9	121	25	10.4
H28581	Hs.92711	ESTs		4.9	108	22	9
N51056	Hs.38891	ESTs		4.9	49	1	4.4
AA430487	Hs.95424	ESTs		4.9	49	7	3.7
AA442868	Hs.13531	ESTs; Weakly similar to (defline not available 5081652) [H		4.9	68	14	1.4
AA435633	Hs.18879	Homo sapiens clone 23965 mRNA sequence		4.9	49	1	4.3
R55185	Hs.3321	ESTs; Highly similar to iroquois-class homeodomain prote		4.9	632	129	1.7
AA257056	Hs.7972	Homo sapiens mRNA for KIAA0871 protein; complete cds		4.9	49	1	4.1
AA455917	Hs.50785	SEC22; vesicle trafficking protein (S. cerevisiae)-like 1		4.9	49	1	4.4
AA329274	Hs.82911	protein tyrosine phosphatase type IVA; member 2		4.9	163	34	15.1
D87969	Hs.82921	CMP-sialic acid transporter		4.9	49	3	3.8
AA451712	Hs.171581	ESTs		4.9	49	1	3.7
X99585	Hs.180139	H.sapiens mRNA for SMT3B protein		4.9	261	53	3.7
M21389	Hs.195850	keratin 5 (epidermolysis bullosa simplex; Dowling-Meara/K		4.8	622	130	0.7
AA001049	Hs.24713	ESTs		4.8	231	49	7.3
AA280670	Hs.24968	ESTs		4.8	79	17	5.2
AA398533	Hs.22209	ESTs		4.8	134	28	3.2
D51095	Hs.35861	ESTs		4.8	48	8	3.1
AA132983	Hs.44155	ESTs; Moderately similar to C-1-TETRAHYDROFOLATE		4.8	96	20	6.5
N69514	Hs.28877	ESTs; Weakly similar to predicted using Genefinder [C.ele		4.8	61	13	5.6
R73468	Hs.140996	ESTs		4.8	48	2	3.4
AA258030	Hs.55356	ESTs; Weakly similar to (defline not available 3874821) [C		4.8	48	1	4.4
AA489046	Hs.94109	ESTs		4.8	179	38	2.8
H88261	Hs.130093	ESTs		4.8	48	1	2.5
H97225	Hs.38592	ESTs		4.8	48	1	0.9
						•	0.0

AA236010	Hs.26613	ESTo	4.0	40		0.5
AA748483	Hs.191356	ESTs Homo sapiens basic transcription factor 2 p44 (btf2p44) g	4.8 4.8	48 48	1 5	0.5 4.1
AA412108	Hs.191803	ESTs	4.8	106	22	1
AA513722	Hs.179729	collagen; type X; alpha 1 (Schmid metaphyseal chondrody	4.8	312	65	30.9
X74987	Hs.12013	ribonuclease L (2';5'-oligoisoadenylate synthetase-depend	4.8	48	8	3.8
AA609427	Hs.210706	ESTs; Moderately similar to !!!! ALU SUBFAMILY SP WAR	4.8	48	1	4.1
AA459555	Hs.31921	Homo sapiens mRNA for KIAA0648 protein; partial cds	4.8	48	1	4.6
W79865	Hs.58367	glypican 4	4.8	48	1	3.6
X06323	Hs.79086	Human MRL3 mRNA for ribosomal protein L3 homologue	4.8	246	51	3.9
AA165231	Hs.8184	ESTs	4.8	53	11	3.7
R38185	Hs.83954	ESTs; Moderately similar to (defline not available 4335943	4.8	153	32	4.3
AA129390	Hs.5285	ESTs CQA1	4.8	93	20	3.1
D14661	Hs.119	gene predicted from cDNA with a complete coding sequen	4.7	119	26	3
D25538	Hs.172199	adenylate cyclase 7	4.7	47	1	4.3
D26361	Hs.3104	KIAA0042 gene product	4.7	47	4	0.7
HG4557-HT4962		"Small Nuclear Ribonucleoprotein U1, 1snrp"	4.7	47	1	4.2
C02582	Hs.109253	ESTs; Highly similar to (defline not available 5114045) [H.	4.7	229	49	7.9
AA040154	Hs.32478	ESTs	4.7	201	43	4.5
AA286809	Hs.28423	ESTs	4.7	152	33	5.3
AA412473	Hs.25880	ESTs	4.7	47	1	4
AA026894	Hs.42826	ESTs; Weakly similar to !!!! ALU CLASS B WARNING EN	4.7	47	4	4.3
N24716	Hs.12244	ESTs	4.7	47	4	4.2
R68425	Hs.28886	ESTs; Moderately similar to !!!! ALU SUBFAMILY SQ WA	4.7	54	12	4.5
AA055768	Hs.122576	ESTs	4.7	770	166	5.8
AA165313	Hs.131189	ESTs	4.7	57	12	4.7
AA478625	110.101100	ESTs	4.7	59	13	4.7
AA044840	Hs.241676	stromal cell-derived factor 1	4.7	114	25	0.9
N90960	Hs.227459	ESTs; Moderately similar to !!!! ALU SUBFAMILY SP WAR	4.7	151	32	9.3
AA873285	Hs.137947	ESTs	4.7	47	3	4.4
D21262	Hs.75337	Human mRNA for KIAA0035 gene; partial cds	4.7	47	1	4
AA214305	Hs.76173	ESTs	4.7	47	1	4.1
AA235803	Hs.9946	ESTs	4.7	710	151	2.5
AA102520	Hs.168017	ESTs; Weakly similar to heat shock protein hsp40 homolo BCH2	4.7	556	119	4.5
AA491465	Hs.28792	ESTs BCU9	4.7	381	81	6.4
AA393803	Hs.16869	ESTs	4.7	747	158	5.7
U02680	Hs.82643	protein tyrosine kinase 9	4.6	148	32	11.3
U18291	Hs.1592	CDC16 (cell division cycle 16; S. cerevisiae; homolog)	4.6	151	33	2
AA476473	Hs.247244	Homo sapiens Trio mRNA; complete cds	4.6	46	1	4
AA609943	Hs.32793	ESTs	4.6	71	16	3.6
T34527	Hs.80120	UDP-N-acetyl-alpha-D-galactosamine:polypeptide N-acet	4.6	199	44	19.2
W19222	Hs.7041	ESTs; Weakly similar to !!!! ALU SUBFAMILY SQ WARNI	4.6	46	4	4.3
AA400247	Hs.42173	ESTs	4.6	46	2	1.8
H29532	Hs.101174	microtubule-associated protein tau	4.6	163	35	7.3
N49408	Hs.136102	Homo sapiens mRNA for KIAA0853 protein; partial cds	4.6	46	1	3.8
W02102	Hs.53565	ESTs; Weakly similar to !!!! ALU SUBFAMILY J WARNING	4.6	60	13	4.8
W69134	Hs.57987	ESTs	4.6	46	7	0.8
AA412488	Hs.48820	ESTs	4.6	46	3	0.8
AA447504	Hs.100261	ESTs	4.6	46	8	3.9
AA437118	Hs.11500	ESTs	4.6	199	44	2.3
AA421139	Hs.173542	ESTs	4.6	239	53	3.5
N64405	Hs.29379	ESTs	4.6	46	7	3.8
AA431459	Hs.47783	ESTs	4.6	46	1	4.3
AA447230	Hs.5070	ESTs	4.6	46	1	4.4
AA135468	Hs.71573	ESTs	4.6	46	5	3.5
R38102	Hs.50421	KIAA0203 gene product	4.6	69	15	5.8
J04177	Hs.82772	collagen; type XI; alpha 1 BCA8	4.6	1216	267	4.4
H25577	Hs.176588	ESTs; Weakly similar to (defline not available 4519535) [H BCB1	4.6	913	199	2.9
AA047036	Hs.62817	ESTs BCR9	4.6	427	93	10.4
AA148885	Hs.154443	minichromosome maintenance deficient (S. cerevisiae) 4	4.6	196	43	10
AA190993	Hs.246174	a disintegrin and metalloproteinase domain 12 (meltrin alp	4.6	132	29	9.7
			-			

D37965	Hs.170040	platelet-derived growth factor receptor-like	4.5	45	4	4
U33052	Hs.69171	protein kinase C-like 2	4.5	45	1	3.6
U71207	Hs.29279	eyes absent (Drosophila) homolog 2	4.5	45	1	2.8
AA195399	Hs.24641	ESTs	4.5	45	1	3.6
AA233168	Hs.3585	ESTs; Weakly similar to coded for by C. elegans cDNA CE	4.5	45	10	0.5
AA281623	Hs.7525	ESTs; Weakly similar to (defline not available 3882205) [H	4.5	45	1	3.7
AA435542	Hs.25796	ESTs	4.5	116	26	4.5
AA489790	Hs.167496	Homo sapiens Ran-GTP binding protein mRNA; partial cd	4.5	45	1	3.8
AA036811	Hs.165030	ESTs	4.5	45	1	3.8
W94942	Hs.177534	ESTs; Weakly similar to dual-specificity protein tyrosine ph	4.5	45	4	2.6
Z40861	Hs.6540	ESTs	4.5	45	2	3
AA253217	Hs.41271	ESTs	4.5	290	65	3.7
AA279943	Hs.122579	ESTs	4.5	96	21	7.8
AA459956	Hs.49163	ESTS	4.5	45	9	3.4
F04816	Hs.92127	ESTs	4.5	96	22	6.9
N39214	Hs.44708	ser-Thr protein kinase related to the myotonic dystrophy p	4.5	211	47	5
AA621785	Hs.170008	ESTs	4.5	45	2	3.6
			4.5	45 45	1	2.4
X56199	Hs.244401	constitutive androstane receptor-beta; orphan nuclear hor	4.5	45 45	1	2.4
Z38919	Hs.21929	ESTs				12
AA430008	Hs.8117	ESTs	4.5	137	31	
D38073	Hs.179565	minichromosome maintenance deficient (S. cerevisiae) 3	4.5	45	2	3.4
U41060	Hs.79136	Human breast cancer; estrogen regulated LIV-1 protein (L B		1472	330	2.1
D78611	Hs.79284	, ,	BC1 4.5	129	29	3.1
T17185	Hs.4299		HA1 4.5	390	87	5.3
U85658	Hs.61796	transcription factor AP-2 gamma (activating enhancer-bind	4.4	255	58	1.6
AA412059	Hs.26864	ESTs	4.4	174	40	1.6
AA452590	Hs.30348	ESTs	4.4	222	51	1.8
AA464708	Hs.249247	ESTs; Weakly similar to alternatively spliced product using	4.4	371	84	3.2
AA009528	Hs.42743	ESTs; Weakly similar to predicted using Genefinder [C.ele	4.4	73	17	6.2
H97678	Hs.31319	ESTs	4.4	103	24	3.8
W92713	Hs.11732	ESTs	4.4	44	6	2.3
Z39211	Hs.150926	fucose-1-phosphate guanylyltransferase	4.4	104	24	5.1
AA024604	Hs.26102	ESTs	4.4	. 44	1	3
AA401474	Hs.208414	ESTs	4.4	44	7	1.1
W72967	Hs.191381	ESTs	4.4	44	1	3.1
AA425887	Hs.98502	ESTs	4.4	48	11	0.9
Al334393	Hs.18113	ESTs	4.4	76	18	1
N59212	Hs.236081	C-terminal binding protein 2	4.4	44	1	3.8
L07493	Hs.1608	replication protein A3 (14kD)	4.4	44	1	4.1
AA281770	Hs.184081	seven in absentia (Drosophila) homolog 1	4.4	53	12	2.1
X54199	Hs.82285	phosphoribosylglycinamide formyltransferase; phosphorib	4.4	44	1	4.1
X94453	Hs.114366	pyrroline-5-carboxylate synthetase (glutamate gamma-sem	4.3	77	18	7.2
AA226968	Hs.22826	ESTs	4.3	43	1	3.9
AA398892	Hs.24391	ESTs	4.3	772	179	1.7
AA399414	Hs.28332	ESTs	4.3	43	1	3.7
AA465093	Hs.239489	TIA1 cytotoxic granule-associated RNA-binding protein	4.3	101	24	1.6
AA489636	Hs.25253	ESTs	4.3	43	10	2.2
AA025728	Hs.61307	ESTs	4.3	43	10	2.7
N59543	Hs.15456	PDZ domain containing 1	4.3	43	1	2
N69113	Hs.110855	ESTs	4.3	43	1	1
N89820	Hs.14559	ESTs	4.3	43	5	2.2
R53439	Hs.194149	ESTs	4.3	45 43	11	4.4
W60439	Hs.119370	ESTs; Moderately similar to cbp146 [M.musculus]	4.3	43	8	3.6
W81552	Hs.242943	constitutive androstane receptor-beta; orphan nuclear hor	4.3	819	191	1.2
N79820	Hs.50854	ESTs	4.3	162	38	12.1
T79274	Hs.10175	ESTs	4.3	65	15	5.7
Z36290	Hs.173933	ESTs; Weakly similar to NUCLEAR FACTOR 1/X [H.sapie	4.3	43	1	3.5
R25607	Hs.23978	scaffold attachment factor B	4.3	68	16	2.8
AA916752 N70516	Hs.244697	ESTs; Highly similar to MEM3 [M.musculus]	4.3	152	35	12.5
N79516	Hs.73287	ESTs	4.3	43	1	3.9

F13665	Hs.65641	ESTs	4.3	190	44	5.4
AB003103	Hs.4295	proteasome (prosome; macropain) 26S subunit; non-ATP	4.2	152	36	12.2
L37936	Hs.3273	Ts translation elongation factor; mitochondrial	4.2	50	12	4.4
U14518	Hs.1594	centromere protein A (17kD)	4.2	42	7	3.4
U90919	Hs.7137	Human clones 23667 and 23775 zinc finger protein mRNA	4.2	42	7	3.7
AA058846	Hs.33363	ESTs	4.2		32	4
AA171736	Hs.35947	ESTs; Highly similar to methyl-CpG binding protein MBD4	4.2		22	2.8
AA227145	Hs.209473	ESTs; Moderately similar to transformation-related protein	4.2		14	2.2
H08778	Hs.133521	ESTs	4.2		14	0.8
R40576	Hs.21590	ESTs; Weakly similar to !!!! ALU SUBFAMILY SC WARNIN	4.2		30	7.4
R66534	Hs.28403	ESTs	4.2		6	3.6
		ESTs	4.2		11	1.4
Z39898	Hs.21948		4.2		9	1.1
AA251524	Hs.44649	ESTs	4.2		1	2.7
F08813	Hs.97413	ret finger protein-like 3 antisense	4.2		30	1.8
AA191353	Hs.22385	ESTs			30 19	1.6
AA412494	Hs.98152	ESTs	4.2			
AA599786	Hs.112110	ESTs	4.2		7	2.9
R01073	Hs.191202	ESTs	4.2		7	3
AA504343	Hs.183475	ESTs; Moderately similar to !!!! ALU SUBFAMILY J WARN	4.2		25	7.8
N30436	Hs.11556	ESTs	4.2		1	3.8
U38847	Hs.151518	TAR (HIV) RNA-binding protein 1	4.2		11	1.1
AA608856	Hs.431	murine leukemia viral (bmi-1) oncogene homolog	4.2		1	2.2
W85888	Hs.47334	ESTs; Moderately similar to !!!! ALU SUBFAMILY SQ WA	4.2		38	7.1
AA453614	Hs.5460	Homo sapiens mRNA for KIAA0776 protein; partial cds	4.2		41	12.6
X57025	Hs.85112	insulin-like growth factor 1 (somatomedin C)	4.2	42	5	2.6
D63391	Hs.6793	platelet-activating factor acetylhydrolase; isoform lb; gamm	4.2	187	44	5.4
AA427861	Hs.59503	ESTs	4.2	79	19	1.9
D14657	Hs.81892	KIAA0101 gene product	4.1	320	78	10.6
D29677	Hs.3085	KIAA0054 gene product	4.1	64	16	3
HG2755-HT2862		T-Plastin	4.1	259	63	1.9
L05425		Homo sapiens autoantigen mRNA; complete cds	4.1	53	13	4
U60808	Hs.152981	CDP-diacylglycerol synthase (phosphatidate cytidylyltrans	4.1	41	1	3.3
U79293	Hs.159264	Human clone 23948 mRNA sequence	4.1	41	1	2.4
X75042	Hs.44313	v-rel avian reticuloendotheliosis viral oncogene homolog	4.1	53	13	4.9
AA236950	Hs.8115	ESTs	4.1	41	2	3.3
H06746	Hs.20072	ESTs	4.1	41	7	1.7
N90430	Hs.6616	ESTs	4.1	41	1	2.6
AA436475	Hs.190104	ESTs	4.1	43	11	1.6
D82808	Hs.17820	Rho-associated; coiled-coil containing protein kinase 1	4.1	41	1	3.6
N77151	Hs.61638	Homo sapiens mRNA for KIAA0799 protein; partial cds	4.1	62	15	4.9
AA093348	Hs.7306	secreted frizzled-related protein 1	4.1	374	91	1.1
W95070	Hs.74316	desmoplakin (DPI; DPII)	4.1	640	158	3
AA243746	Hs.211577	ESTs; Highly similar to CG1 protein [H.sapiens]	4.1	301	73	6.1
AA169379	Hs.72865		:U4 <b>4.</b> 1	334	82	3.4
AA490890	Hs.105273	ESTs	4.1	72	18	1.5
D86961	Hs.79299	lipoma HMGIC fusion partner-like 2	4	40	1	3.8
L28997	Hs.77102	ADP-ribosylation factor-like 1	4	110	28	10.7
AA281245	Hs.23317	ESTs	4	75	19	1.7
AA393793	Hs.110347	ESTs; Highly similar to (defline not available 4468913) [H.	4	40	3	3.2
AA171755	Hs.181915	ESTs	4	40	7	1.1
R51818	Hs.104222	ESTs	4	70	18	6.8
W72471	Hs.23920	ESTs	4	48	12	4
T23820	Hs.155478	cyclin T2	4	40	4	1.2
T56679	Hs.865	RAP1A; member of RAS oncogene family	4	40	1	3.4
H03686	Hs.220689	Ras-GTPase-activating protein SH3-domain-binding prote	4	40	4	3.2
N78483	Hs.24809	ESTs	4	95	24	1.1
U90551	Hs.28777	H2A histone family; member L	4	350	88	3
AA010163	Hs.3383	upstream regulatory element binding protein 1	4		35	1.8
W60186	Hs.169487	ESTs	4		114	2
L04656	113.103707	carbonic anhydrase VIII	3.9		8	3.6
204000		ouround annyarado vin	J.,	. 39	J	3.0

		A A TOTAL A CONTRACT	• •		_	2.5
M27492	Hs.82112	interleukin 1 receptor; type I	3.9	39	2 11	3.5 1
X87613	Hs.169344	H.sapiens mRNA for skeletal muscle abundant protein	3.9 3.9	43 49	13	2.5
Y09443	Hs.22580	alkylglycerone phosphate synthase			6	3.2
AA227448	Hs.5003	Homo sapiens mRNA for KIAA0456 protein; partial cds	3.9	39 70		
AA235303	Hs.8645	ESTs	3.9	79	20	6.5
AA398197	Hs.30029	ESTs	3.9	371	94	4.6
AA609210	Hs.19221	ESTs	3.9	98	25	8.6
N36001	Hs.17348	ESTs; Weakly similar to alternatively spliced product using	3.9	353	90	1.2
N67437	Hs.24375	ESTs	3.9	146	37	9.8
AA427528	Hs.114547	ESTs; Weakly similar to ZINC FINGER PROTEIN 84 [H.sa	3.9	39	8	2.2
AA521080	Hs.46765	ESTs	3.9	39	10	0.6
AA255933	Hs.109111	ESTs	3.9	161	42	2
AA400412	Hs.97794	ESTs	3.9	39	1	0.2
AA425374	Hs.193063	ESTs	3.9	39	1	0.2
AA600121	Hs.190253	ESTs	3.9	39	1	3.2
AA609471	Hs.112712	ESTs	3.9	39	5	3.7
AA370120	Hs.7870	ESTs; Weakly similar to Ylr350wp [S.cerevisiae]	3.9	47	12	4.4
H02682	Hs.99189	ESTs; Weakly similar to novel stromal cell protein [M.mus	3.9	220	57	2.5
M28879	Hs.1051	granzyme B (granzyme 2; cytotoxic T-lymphocyte-associa	3.9	43	11	1.8
D13435	Hs.166982	phosphatidylinositol glycan; class F	3.9	54	14	5.1
N21679	Hs.180059	ESTs	3.9	39	1	1.9
D31263	Hs.15929	ESTs	3.9	39	1	2.6
AA477739	Hs.246856	ESTs	3.9	39	1	3.4
R49035	Hs.26176	ESTs	3.9	116	30	0.5
U66615	Hs.172280	SWI/SNF related; matrix associated; actin dependent regu	3.9	39	1	2.5
U23070	Hs.78776	Human putative transmembrane protein (nma) mRNA; co BCH9	3.9	442	114	1.3
N22107	Hs.172241	ESTs; Moderately similar to !!!! ALU SUBFAMILY SC WARBCN7	3.9	322	83	4.4
AA609651	Hs.112742	ESTs BCX7	3.9	60	16	4.8
K01160		Accession not listed in Genbank	3.9	390	100	11.1
AA057193	Hs.25252	ESTs	3.9	280	72	3.3
D28137	Hs.118110	bone marrow stromal cell antigen 2	3.8	350	93	1.9
J05249	Hs.79411	replication protein A2 (32kD)	3.8	115	30	7.1
S80562	Hs.194662	calponin 3; acidic	3.8	399	105	3.3
U28368	Hs.34853	inhibitor of DNA binding 4; dominant negative helix-loop-h	3.8	163	43	0.5
U57721	Hs.81771	kynureninase; I-kynurenine hydrolase	3.8	38	1	1.5
Z74615	Hs.172928	collagen; type I; alpha 1	3.8	1612	429	3.1
R86920	Hs.127585	ESTs	3.8	38	4	1.2
AA027317	Hs.221929	ESTs; Weakly similar to !!!! ALU SUBFAMILY J WARNING	3.8	40	11	3.8
AA084602	Hs.29669	ESTs	3.8	38	1	2.4
AA179826	Hs.32058	ESTs	3.8	38	2	3.2
AA233790	Hs.4104	ESTs	3.8	93	25	7.5
AA424006	Hs.22972	ESTs; Weakly similar to steroid 5-alpha-reductase 2 [H.sa	3.8	38	1	3.3
AA429951	Hs.21104	ESTs	3.8	83	22	7.5
AA436836	Hs.35580	ESTs	3.8	38	1	1.9
AA456646	Hs.28661	ESTs	3.8	263	69	3.9
AA489009	Hs.26994	ESTs	3.8	38	1	1.6
AA079468	Hs.94631	ESTs	3.8	38	1	3.2
AA179387	Hs.25264	ESTs	3.8	233	62	3.8
H42396	Hs.107872	ESTs	3.8	38	7	2.8
Z38909	Hs.22265	ESTs	3.8	73	19	1.8
AA478729	Hs.76450	ESTs	3.8	38	7	2.1
AA347422	Hs.238040	ESTs; Weakly similar to hypothetical protein [H.sapiens]	3.8	38	7	0.2
AA485458	Hs.181357	ESTs; Moderately similar to laminin-binding protein [H.sap	3.8	207	55	5.5
H05323	Hs.247486	ESTs	3.8	58	16	5.5
AA512902	Hs.7337	ESTs	3.8	38	1	3
N75007	Hs.199009	ESTs; Weakly similar to (defline not available 4589652) [H	3.8	38	1	0.9
AA232276	Hs.22806	ESTs; Weakly similar to !!!! ALU SUBFAMILY J WARNING	3.8	42	11	0.6
AA465527	Hs.23853	ESTs	3.8	38	1	3.3
AA418039	Hs.26155	ESTs	3.8	38	1	3
AA262821	Hs.28578	ESTs	3.8	79	21	6.9

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AA599653	Hs.30696	transcription factor-like 5 (basic helix-loop-helix)		3.8	51	14	1.7
AA448297	Hs.237825	calcium/calmodulin-dependent protein kinase (CaM kinas		3.8	38	1	3
AA174183	Hs.93872	ESTs		3.8	240	64	3.2
C00038	Hs.23579	ESTs	BCQ5	3.8	585	153	3.7
N91023	Hs.170057	ESTs	BCW4	3.8	425	111	4
AA007160	Hs.14846	ESTs	CQA2	3.8	82	22	3.1
Al167942	Hs.61635	Homo sapiens BAC clone RG041D11 from 7q21	PAA5	3.8	38	1	2.7
AA569531	Hs.162859	ESTs	PAA6	3.8	38	7	3.4
HG4297-HT4567	113.102003	Transcriptional Coactivator Pc4		3.7	477	130	3.1
X53961	Hs.347	lactotransferrin		3.7	1421	388	1.9
AA434508	113.547	"zw31c1.r1 Soares ovary tumor NbHOT Homo sapiens ct	)	3.7	37	7	2.1
R64534	Hs.101469	ESTs		3.7	37	5	2.5
AA126855	Hs.13268	ESTs		3.7	157	43	3.6
AA128548	Hs.90847	ESTs; Weakly similar to Similarity with yeast transcription		3.7	37	1	3.2
H03627	Hs.245209	ESTs		3.7	37	4	2
H53572	Hs.32407	ESTs		3.7	37	1	2.1
	Hs.15119	ESTs		3.7	119	33	6.7
N68869	Hs.25978	ESTs		3.7	37	1	3
R52949		ESTs; Moderately similar to FK506-binding protein 65kD		3.7	239	65	3.6
W80763	Hs.3849	ESTs. Woderately similar to 1 1000-binding protein cons	1	3.7	37	1	1.8
AA504116	Hs.82501	H.sapiens mRNA similar to Xenopus laevis mRNA for KD	F	3.7	37	6	0.5
N94475	Hs.227342		<b></b>	3.7	37	4	3
R46061	Hs.92482	ESTs ESTs; Weakly similar to !!!! ALU SUBFAMILY J WARNIN	c.	3.7	37	4	1.9
W84767	Hs.58698	•	0	3.7	590	159	3.8
W86835	Hs.14158	copine III		3.7	37	1	0.5
AA280738	Hs.128679	ESTs		3.7	37	10	1
AA399441	Hs.104699	ESTs		3.7	41	11	1.6
AA495812	Hs.105364	ESTs		3.7	39	11	3.2
R28587	Hs.11000	Homo sapiens brain my047 protein mRNA; complete cds		3.7	37	1	3.4
M87339	Hs.35120	replication factor C (activator 1) 4 (37kD)		3.7	37	9	2.8
F09788	Hs.3622	procollagen-proline; 2-oxoglutarate 4-dioxygenase (prolin		3.7	37	1	2.2
X75535	Hs.168670	peroxisomal farnesylated protein	_	3.7	91	25	2.6
D15050	Hs.232068	"Human mRNA for transcription factor AREB6, complete	U	3.7	41	11	0.6
W58612	Hs.173840	ESTs	BCU1	3.7	625	168	3.8
AA121315	Hs.70823	KIAA1077 protein	BÇU I	3.7	162	44	12.4
AA477445	Hs.105341	ESTs		3.7	224	61	21
AA477571	Hs.152601	UDP-glucose ceramide glucosyltransferase Human mRNA for KIAA0225 gene; partial cds		3.6	36	7	3.2
D86978	Hs.84790			3.6	36	1	2.6
M55542	Hs.62661	guanylate binding protein 1; interferon-inducible; 67kD carboxypeptidase B1 (tissue)		3.6	824	227	1.4
M81057	Hs.180884			3.6	142	39	1.6
U90304	Hs.25351	iroquois-class homeodomain protein ESTs		3.6	36	1	0.1
AA282138	Hs.11325	ESTs		3.6	68	19	6
AA398346	Hs.21898 Hs.23505	ESTs		3.6	36	7	3.1
AA399623		ESTs; Moderately similar to UDP-GLUCOSE:GLYCOPR	2	3.6	45	13	1.3
AA400517	Hs.22983	C-terminal binding protein 2	•	3.6	444	125	4.6
AA417287	Hs.171391	Homo sapiens clone 24416 mRNA sequence		3.6	365	103	6.9
AA417761 AA461495	Hs.5957 Hs.14512	ESTs		3.6	210	58	4.7
AA489665	Hs.25245	ESTs		3.6	36	1	1.2
D59368	Hs.159872	ESTs		3.6	36	6	0.5
H53829	Hs.36823	ESTs		3.6	36	10	2.5
T93630	Hs.17207	ESTs		3.6	36	4	2.6
AA456020	Hs.50848	ESTs; Weakly similar to (defline not available 4239895)	н	3.6	36	1	2
F01601	Hs.117485	ESTs		3.6	36	1	1.9
H99959	Hs.42768	ESTs; Weakly similar to (defline not available 4689264)	Н	3.6	41	12	2.8
N66413	Hs.172466	ESTs; Weakly similar to (defline not available 3882271)		3.6	89	25	0.9
W73788	Hs.43213	ESTs	•	3.6	36	1	2.9
AA280794	Hs.241328	ESTs		3.6	36	8	0.2
AA426270	Hs.145696	ESTs		3.6	150	42	3.2
AA465196	Hs.107233	ESTs		3.6	36	1	3.4
W38240		Accession not listed in Genbank		3.6	38	11	2.6

				2.6	36	6	2.9
AA714635	Hs.181297	ESTs "EST176522 Colon carcinoma (Caco-2) cell line II Homo s		3.6 3.6	121	34	11.8
AA305536 AA129640	Hs.128065	ESTs		3.6	36	10	1.9
D00763	Hs.243746	proteasome (prosome; macropain) subunit; alpha type; 4		3.6	130	36	3.5
D86959	Hs.105751	KIAA0204 gene product		3.6	36	5	1.5
X55330	Hs.207776	aspartylglucosaminidase		3.6	36	1	2.7
M95767	Hs.135578	chitobiase; di-N-acetyl-		3.6	36	1	1.2
	Hs.19347	ESTs		3.6	100	28	6.6
AA248406	Hs.246093	ESTs		3.6	141	39	12.6
AA234767	Hs.46967	ESTs		3.6	36	1	3.1
AA479933		ESTs; Weakly similar to ZINC FINGER PROTEIN 135 [H.		3.6	146	41	1.1
AA287107	Hs.172945 Hs.64311	a disintegrin and metalloproteinase domain 17 (tumor nec		3.6	36	1	3.1
U69611	Hs.194369	Homo sapiens chromosome 1 atrophin-1 related protein (		3.6	39	11	0.4
AA284143	Hs.166975	Human splicing factor SRp40-1 (SRp40) mRNA; complete	1	3.6	36	1	0.4
U30827	Hs.250165	Lon protease-like protein		3.6	36	1	2.8
X76040 N68921	Hs.34806	ESTs; Weakly similar to neogenin [H.sapiens]	BCW3	3.6	402	112	4.9
T95333	Hs.122730	ESTs; Weakly similar to Strabismus [D.melanogaster]	CVA9	3.6	224	63	4
H95039	Hs.32168	ESTs		3.6	282	79	1.7
AA609710	Hs.42582	ESTs; Weakly similar to similar to GTP-binding protein [C.	_	3.6	256	72	3.7
D42084	Hs.82007	Human mRNA for KIAA0094 gene; partial cds		3.5	96	28	1.3
	Hs.75909	Human mRNA for KIAA0182 gene; partial cds		3.5	78	23	4.8
D80004	Hs.122669	Human mRNA for KIAA0264 gene; partial cds		3.5	35	9	3.1
D87453	115.122005	"Mucin 1, Epithelial, Alt. Splice 6"		3.5	37	11	2.8
HG371-HT1063	Hs.151461	embryonic ectoderm development protein		3.5	35	1	2.7
U90651	Hs.14600	ESTs		3.5	35	7	2.7
AA232215	Hs.25242	ESTs		3.5	73	21	1.6
AA258873	Hs.23019	ESTs; Weakly similar to ZINC FINGER PROTEIN 135 [H.		3.5	53	15	1.2
AA417034 AA482035	Hs.28070	KIAA0753 gene product		3.5	58	17	1.6
AA504144	Hs.22315	ESTs		3.5	180	52	2.3
T74445	Hs.5957	"yc82f8.r1 Soares infant brain 1NIB Homo sapiens cDNA		3.5	35	1	2.6
AA016021	Hs.173091	Homo sapiens HCG-1 protein (HCG-1) mRNA; complete		3.5	282	80	3.7
AA025315	Hs.61184	ESTs		3.5	35	1	1
AA129968	Hs.49376	ESTs; Weakly similar to PROTEIN PHOSPHATASE PP2	Α	3.5	35	1	3.2
H89315	110.10070	"yw25e09.s1 Morton Fetal Cochlea Homo sapiens cDNA		3.5	35	8	1.9
N51374	Hs.96870	ESTs		3.5	35	9	3.2
R08850	Hs.9786	ESTs		3.5	35	1	2.1
R33468	Hs.24651	ESTs		3.5	105	30	9.6
R49482	Hs.5637	ESTs		3.5	507	145	3.3
R49483	Hs.22159	ESTs; Weakly similar to finger protein HZF10; Krueppel-r	e	3.5	40	12	2.5
R54822	Hs.26244	ESTs		3.5	41	12	3.7
T66847	Hs.194040	ESTs		3.5	35	1	1.4
W96222	Hs.34192	ESTs		3.5	35	6	3.2
AA459703	Hs.79070	ESTs; Moderately similar to coded for by C. elegans cDN	IA	3.5	35	8	3.3
N24954	Hs.42502	ESTs		3.5	72	21	1.3
N89881	Hs.44577	ESTs		3.5	35	1	2.9
AA377296	Hs.97104	ESTs		3.5	37	11	0.1
AA410383	Hs.100431	B-cell-homing chemokine (ligand for Burkitt's lymphoma	re	3.5	35	1	2.6
AA412151	Hs.235402	ESTs		3.5	143	41	2.6
AA428213	Hs.98523	ESTs		3.5	35	3	2.3
AA454103	Hs.110031	ESTs		3.5	35	1	3
Al479264	Hs.13058	ESTs		3.5	35	4	3.3
AA936428	Hs.128638	ESTs		3.5	35	1	3.1
A1369384		arylsulfatase D		3.5	113	33	1.7
U58522	Hs.155485	"Human huntingtin interacting protein (HIP2) mRNA, com	ηp	3.5	79	23	2.5
U25997	Hs.25590	stanniocalcin		3.5	402	114	2.1
AA090617	Hs.247614	ESTs		3.5	35	1	2.5
AA599801	Hs.239507	ESTs		3.5	73	21	6.3
AA496037	Hs.60293	ESTs		3.5	110	32	2.1
N30704	Hs.238797	ESTs	DC 4.0	3.5	35 706	7	2.1
X72755	Hs.77367	monokine induced by gamma interferon	BCA6	3.5	796	228	3.2

W72838	Hs.58213	ESTs	BCH1	3.5	2073	595	2.1
R45698	Hs.21893	ESTs	CZA1	3.5	124	35	6.5
X54941	Hs.77550	CDC28 protein kinase 1		3.5	332	94	3.1
Y00815	Hs.75216	protein tyrosine phosphatase; receptor type; F		3.5	564	162	1.7
D31888	Hs.78398	Homo sapiens clone 24709 mRNA sequence		3.4	77	23	5.9
D87470	Hs.75400	Human mRNA for KIAA0280 gene; partial cds		3.4	34	1	1.2
X85750	Hs.79889	H.sapiens mRNA for transcript associated with monocyte		3.4	34	8	2.3
AA004211	Hs.30977	ESTs; Weakly similar to putative p150 [H.sapiens]		3.4	154	46	3
AA292711	Hs.29131	ESTs		3.4	34	1	3.1
AA400093	Hs.32271	ESTs; Weakly similar to !!!! ALU SUBFAMILY SP WARNII	V	3.4	34	8	2.9
AA401633	Hs.22380	ESTs		3.4	34	1	1.5
AA412505	Hs.10653	ESTs		3.4	49	15	4.4
AA417067	Hs.13055	ESTs		3.4	116	35	2.2
D51235	Hs.82689	tumor rejection antigen (gp96) 1		3.4	251	74	23.7
AA044181	Hs.62677	ESTs		3.4	34	1	2.3
AA129933	Hs.71168	Homo sapiens clone 24674 mRNA sequence		3.4	34	1	2.8
AA156460	Hs.44229	ESTs		3.4	34	1	2.6
AA167708	Hs.52184	ESTs		3.4	71	21	2.4
N48603	Hs.14947	ESTs		3.4	115	34	2.4
N95837	Hs.169111	ESTs		3.4	314	91	2.4
AA456968	Hs.82669	ESTs		3.4	34	8	1
N79496	Hs.50824	EST		3.4	740	217	2.8
T78324	Hs.90905	ESTs		3.4	34	3	2.4
W73057	Hs.58272	ESTs; Moderately similar to alternatively spliced product u	ı	3.4	34	1	2.5
AA171739	Hs.101590	ESTs		3.4	34	1	1.7
AA251973	Hs.143853	ESTs		3.4	34	4	0.1
AA406293	Hs.193498	ESTs		3.4	34	1	0.1
AA418988	Hs.98314	ESTs		3.4	34	10	0.7
AA598899	Hs.112493	ESTs		3.4	34	10	2.6
AA621348	Hs.227933	ESTs; Highly similar to (defline not available 5281121) [H.		3.4	80	24	3.8
R41933	Hs.140237	ESTs; Weakly similar to neuronal thread protein AD7c-NT		3.4	210	63	3.3
T91518	113.140231	"ye20f05.s1 Stratagene lung (#937210) Homo sapiens cD		3.4	985	286	2.8
R56892	Hs.75544	tyrosine 3-monooxygenase/tryptophan 5-monooxygenase		3.4	234	68	10.7
AA219081	Hs.242396	ESTs; Moderately similar to !!!! ALU SUBFAMILY SB2 WA		3.4	107	32	9.9
W46810	Hs.20521	HMT1 (hnRNP methyltransferase; S. cerevisiae)-like 2	•	3.4	525	154	5.3
X86098	Hs.3238	adenovirus 5 E1A binding protein		3.4	115	34	9.1
H96226	Hs.42151	ESTs		3.4	58	17	4
N41849	Hs.119410	Homo sapiens cytokine receptor related protein 4 (CYTOF	2	3.4	34	2	3
AA386264	Hs.5337	ESTs	`	3.4	193	58	12.3
X78627	Hs.75066	translin		3.4	178	53	8.8
C02374	Hs.7822	ESTs		3.4	91	27	8.5
C06270	Hs.8078	Homo sapiens clone FBD3 Cri-du-chat critical region mRN	J	3.4	34	5	2.6
AA481414	Hs.8868	golgi SNAP receptor complex member 1	•	3.4	179	52	1.5
AA262727	Hs.12144	ESTs		3.4	88	26	1.4
M77698	Hs.97496	YY1 transcription factor		3.4	475	142	2.5
AA215333	Hs.97101	ESTs	CXA1	3.4	169	50	9.1
X70683	Hs.83484	SRY (sex determining region Y)-box 4		3.4	496	144	1.6
N81017	Hs.42679	ESTs		3.4	118	35	2.3
HG2874-HT3018		Ribosomal Protein L39 Homolog		3.3	116	36	2.2
HG4036-HT4306		Retinoblastoma 1		3.3	33	1	0.8
M84605	Hs.957	Human putative opioid receptor mRNA; complete cds		3.3	36	11	2.4
U43286	Hs.118725	Human selenophosphate synthetase 2 (SPS2) mRNA; co		3.3	111	34	7.5
X68733	Hs.234726	alpha-1-antichymotrypsin		3.3	1497	458	2.1
Z35402	Hs.194657	"H.sapiens gene encoding E-cadherin, exon 3 and joined		3.3	745	229	1.8
AA251297	Hs.23439	ESTs		3.3	206	63	2.2
AA350771	Hs.17850	ESTs		3.3	98	30	4.7
AA427816	Hs.11803	ESTs		3.3	95	29	4.4
AA434441	Hs.173859	frizzled (Drosophila) homolog 7		3.3	97	30	6.4
AA487561	Hs.5566	ESTs; Highly similar to RAS-RELATED PROTEIN RAB-1/	A	3.3	696	214	1.8
AA598820	Hs.3530	TLS-associated serine-arginine protein		3.3	228	69	2.8

AA137078	Hs.173648	ESTs	3.3	33	5	1.7
AA171529	Hs.183887	ESTs	3.3	33	1	2.9
AA227119	Hs.171558	sex comb on midleg (Drosophila)-like 2	3.3	39	12	1.5
H61560	Hs.161011	EST	3.3	33	1	1.8
H98653	Hs.188006	Homo sapiens mRNA for KIAA0878 protein; complete cds	3.3	138	42	3.6
N69287	Hs.21943	ESTs	3.3	33	1	1.1
R44538	Hs.140889	ESTs	3.3	33	10	2.3
W37382	Hs.11090	ESTs	3.3	744	227	2.5
W42845	Hs.14611	dual specificity phosphatase 11 (RNA/RNP complex 1-inte	3.3	180	54	2.1
Z39742	Hs.247047	ESTs	3.3	33	1	2.4
AA150043	Hs.11498	ESTs	3.3	168	51	7.3
AA261819	Hs.88367	ESTs	3.3	33	1	3
AA481256	Hs.88201	ESTs; Weakly similar to (defline not available 3859560) [H	3.3	106	33	9.8
D51276	Hs.81915	leukemia-associated phosphoprotein p18 (stathmin)	3.3	931	279	5.6
H91164	Hs.237404	ESTs; Moderately similar to !!!! ALU SUBFAMILY SQ WA	3.3	33	1	2.3
N67889	Hs.49397	ESTs	3.3	81	25	1.5
N98488	Hs.161545	EST	3.3	36	11	0.6
R39261	Hs.90790	ESTs	3.3	167	51	2.6
W43000	Hs.159225	ESTs	3.3	33	8	0.9
W52480	Hs.56148	ESTs; Moderately similar to (defline not available 5360125	3.3	33	10	0.5
AA227837	Hs.210566	ESTs; Moderately similar to !!!! ALU SUBFAMILY SC WAR	3.3	33	10	2.8
AA347419	Hs.96870	ESTs	3.3	33	3	0.2
AA446190	Hs.99051	ESTs	3.3	53	16	4
AA480103	Hs.111730	ESTs	3.3	207	63	3.5
N53976	Hs.179864	ESTs	3.3	33	1	1.7
W72949	Hs.77495	Human mRNA for KIAA0242 gene; partial cds	3.3	34	11	3.2
H22147	Hs.245474	ESTs; Moderately similar to !!!! ALU SUBFAMILY SP WAR	3.3	33	9	0.9
U31875	Hs.152677	short-chain alcohol dehydrogenase family member	3.3	105	32	3
W15528	Hs.106390	ESTs	3.3	288	87	7.9
X84373	Hs.155017	nuclear receptor interacting protein 1	3.3	354	108	4
X60486	Hs.46423	H4 histone family; member G	3.3	979	298	2.2
AA176475	Hs.4864	Homo sapiens mRNA for KIAA0892 protein; partial cds	3.3	33	4	2.9
AA253330	Hs.5344	ESTs	3.3	909	274	3.2
N26645	Hs.58220	Homo sapiens clone 24723 mRNA sequence	3.3	61	19	5.1
AA142857	Hs.234896	ESTs; Highly similar to geminin [H.sapiens]	3.3	106	33	2.6
AA216562	Hs.69855	neuroblastoma RAS viral (v-ras) oncogene homolog	3.3	33	1	2.9
M33882	Hs.76391	myxovirus (influenza) resistance 1; homolog of murine (in	3.3	380	114	4.9
W84712	Hs.7753	calumenin	3.3	889	267	5
S80437	Hs.83190	"fatty acid synthase {3' region} [human, breast and HepG2	3.3	710	217	2
N21407	Hs.247471	ESTs	3.3	33	1	2
C15324	Hs.93668	ESTs	3.3	1296	394	2.2
D30756	Hs.244822	membrane component; chromosome 17; surface marker 2	3.3	33	1	2.6
T32108	Hs.153315	ESTs CHA4	3.3	571	171	2.0
AA456687	Hs.26057	ESTs	3.3	775	233	2.4
D86969	Hs.82292	KIAA0215 gene product	3.2	32	2	2.9
X06272	Hs.75730	signal recognition particle receptor ('docking protein')	3.2	58	18	5
X15875	Hs.198166	activating transcription factor 2	3.2	32	4	2.6
Z35491	Hs.41714	BCL2-associated athanogene	3.2	41	13	2.8
Z80781	Hs.249216	H2B histone family; member J	3.2	32	5	2.8
AA314389	Hs.203994	ESTs; Weakly similar to (defline not available 4502227) [H	3.2	32	9	2.7
D55869	Hs.173138	ESTs	3.2	32	7	2.4
AA088228	Hs.18272	ESTs	3.2	522	165	1.9
AA112361	Hs.10592	ESTs	3.2	32	4	2.9
AA148859	Hs.179909	ESTs; Moderately similar to !!!! ALU SUBFAMILY J WARN	3.2	32	1	3
AA255874	Hs.23458	ESTs	3.2	466	146	8.4
AA256996	Hs.3862	ESTs	3.2	32	6	1.5
AA279991	Hs.124691	ESTs	3.2	32	1	1
AA369245	Hs.17448	ESTs; Moderately similar to !!!! ALU SUBFAMILY SC WAR	3.2	143	46	3.6
AA419609	Hs.170121	ESTs; Weakly similar to ETX1 {alternatively spliced} [H.sa	3.2	267	83	2.3
AA436628	Hs.158249	KIAA0406 gene product	3.2	37	12	2.6
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T00474		50T M. J				
T63174	Hs.193700	ESTs; Moderately similar to !!!! ALU SUBFAMILY SB WAR	3.2	110	35	9.6
AA004806	Hs.60090	ESTs	3.2	32	5	2.1
AA111879	Hs.69507	EST	3.2	32	5	1.7
AA180453	Hs.73643	ESTs	3.2	32	1	2.2
AA233342	Hs.90680	ESTs; Weakly similar to Unknown gene product [H.sapien	3.2	, 286	91	5.7
F10024	Hs.220640	ESTs	3.2	41	13	3.3
H09594	Hs.10299	ESTs; Moderately similar to !!!! ALU SUBFAMILY SQ WA	3.2	136	43	3.6
R38436	Hs.21181	ESTs	3.2	37	12	3.5
T23860	Hs.7312	ESTs	3.2	99	31	3.1
W60002	Hs.4114	plastin 3 (T isoform)	3.2	238	75	2.1
AA258116	Hs.191533	ESTs; Weakly similar to !!!! ALU SUBFAMILY SQ WARNI	3.2	49	16	4.2
AA435946	Hs.50831	ESTs	3.2	40	13	0.7
AA496000	Hs.4084	ESTs	3.2	173	55	3
N30205	Hs.93740	ESTs; Moderately similar to !!!! ALU SUBFAMILY SX WAR	3.2	35	11	0.7
N66763	Hs.43080	ESTs	3.2	378	117	2.8
Z41050	Hs.108787	Homo sapiens Mcd4p homolog mRNA; complete cds	3.2	106	34	3.3
AA279654	Hs.194524	ESTs; Moderately similar to !!!! ALU SUBFAMILY SX WAR	3.2	32	5	0.6
AA287389	Hs.98267	ESTs	3.2	32	8	3
AA416568	Hs.98203	ESTs	3.2	32	1	0.8
AA431751	Hs.106711	eukaryotic translation initiation factor 4E binding protein 3	3.2	88	28	1.2
AA449121	Hs.99210	ESTs	3.2	291	91	4
AA454149	Hs.99357	EST	3.2	32	10	3.1
AA460324	Hs.99527	ESTs	3.2	36	11	2.5
H90150	Hs.13366	Homo sapiens mRNA; cDNA DKFZp566F133 (from clone	3.2	32	1	1.5
W78134	Hs.122647	N-myristoyltransferase 2	3.2	37	12	3.6
AA488132	Hs.62741	ESTs	3.2	32	10	
L12350	Hs.108623	thrombospondin 2	3.2			2.9
		•		814	257	2.4
J04076	Hs.1395	early growth response 2 (Krox-20 (Drosophila) homolog)	3.2	32	1	0.2
D80074	Hs.169833	ESTs; Highly similar to (defline not available 4689144) [H.	3.2	87	27	1.7
H44386	Hs.22666	ESTs	3.2	210	66	3.8
U26174	Hs.3066	granzyme K (serine protease; granzyme 3; tryptase II)	3.2	82	26	6.6
N51260	Hs.196275	ESTs	3.2	207	64	5.5
X52947	Hs.74471	gap junction protein; alpha 1; 43kD (connexin 43)	3.2	351	111	5.2
Y00264	Hs.177486	amyloid beta (A4) precursor protein (protease nexin-II; Alz	3.2	226	71	2.8
L08044	Hs.169224	trefoil factor 3 (intestinal)	3.2	1872	592	3.3
X04011	Hs.88974	cytochrome b-245; beta polypeptide (chronic granulomato	3.2	143	45	13.9
X87241	Hs.166994	FAT tumor suppressor (Drosophila) homolog	3.2	153	48	4.7
L14922	Hs.166563	replication factor C (activator 1) 1 (145kD)	3.2	32	1	2.4
M23263	Hs.99915	androgen receptor (dihydrotestosterone receptor; testicula	3.2	117	37	9.4
AA156897	Hs.72157	ESTs; Highly similar to (defline not available 4884194) [H. BCU3	3.2	725	227	3.2
AA521354	Hs.24758	ESTs BCW1		266	83	1.8
D21255	Hs.75929	cadherin 11 (OB-cadherin; osteoblast)	3.2	560	174	2.6
AA404352	Hs.178603	ESTs	3.2	372	115	2.1
D43772	Hs.86859	growth factor receptor-bound protein 7	3.1	306	98	1.5
L20320	Hs.184298	cyclin-dependent kinase 7 (homolog of Xenopus MO15 cd	3.1	118	38	2
M55998		"Human alpha-1 collagen type I gene, 3' end"	3.1	2898	923	2.2
M93036	Hs.692	membrane component; chromosomal 4; surface marker (3	3.1	353	116	2.8
U27185	Hs.32943	retinoic acid receptor responder (tazarotene induced) 1	3.1	31	1	1.3
AA148516	Hs.35156	ESTs; Moderately similar to !!!! ALU SUBFAMILY J WARN	3.1	31	1	2.5
AA447223	Hs.25320	ESTs	3.1	72	23	5
AA448850	Hs.17138	ESTs	3.1	165	54	1.6
AA449741	Hs.4029	Glioma-amplified sequence-41	3.1	31	1	2.6
AA599472	Hs.247309	eukaryotic translation elongation factor 1 delta (guanine n	3.1	55	18	3.8
AA600310	Hs.18720	ESTs; Highly similar to (defline not available 4323587) [H.	3.1	75	24	2.2
AA609053	Hs.35198	ESTs	3.1	367	119	2.3
AA025782	Hs.61284	ESTs	3.1	31	9	2.2
AA135894	Hs.194691	retinoic acid induced 3	3.1	529	170	4.1
N29454	Hs.27552	ESTs; Weakly similar to putative p150 [H.sapiens]	3.1	31	1	2.7
N40981	Hs.9856	ESTs	3.1	31	1	1.3
T92735	Hs.17061	ESTs	3.1	453	148	7
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Z38239	Hs.26962	ESTs		3.1	31	5	1.5
AA417375	Hs.76917	ESTs; Weakly similar to KIAA0522 protein [H.sapiens]		3.1	58	19	2.5
AA620761	Hs.47274	Homo sapiens mRNA; cDNA DKFZp564B176 (from clone		3.1	119	39	2
C20652	Hs.68501	ESTs		3.1	31	4	1.9
H95785	Hs.167652	ESTs; Highly similar to CENP-E protein [H.sapiens]		3.1	38	13	1.7
H98153	Hs.42500	ESTs		3.1	295	96	27.9
N49967	Hs.46624	ESTs		3.1	31	1	2.7
N66845	Hs.165411	ESTs; Weakly similar to !!!! ALU CLASS B WARNING EN		3.1	199	64	1
AA382275	Hs.97128	ESTs		3.1	31	1	0.4
AA436890	Hs.98918	ESTs		3.1	31	1	1
AA449453	Hs.192915	ESTs; Weakly similar to !!!! ALU SUBFAMILY J WARNING		3.1	31	6	0.8
AA608588	Hs.193634	ESTs		3.1	927	295	2.1
H88296	Hs.5123	ESTs; Weakly similar to (defline not available 4960208) [H		3.1	41	14	2.7
N50641	Hs.80285	ESTs		3.1	31	1	1.8
T90297	Hs.140571	ESTs		3.1	31	6	2.4
W42680	Hs.95941	ESTs		3.1	31	1	2.8
AA643322	110.00011	"nr59c06.s1 NCI_CGAP_Lym3 Homo sapiens cDNA clon		3.1	31	1	2.5
AA625690	Hs.190272	ESTs		3.1	33	11	2.3
AI354332	Hs.72365	ESTs		3.1	31	8	2
AA093378	Hs.101810	ESTs; Weakly similar to !!!! ALU SUBFAMILY SC WARNIN	1	3.1	172	55	3.1
R48943	Hs.10315	solute carrier family 7 (cationic amino acid transporter; y+	•	3.1	31	1	2.2
W93562	Hs.105749	Homo sapiens mRNA for KIAA0553 protein; partial cds		3.1	34	11	2.7
		ESTs		3.1	31	2	2.5
F11087	Hs.239666	ESTs		3.1	31	3	3
T71333	Hs.13854	X-box binding protein 1		3.1	1336	434	1.4
M31627	Hs.149923	ESTs		3.1	197	63	18.7
AA121127	Hs.181307	ESTs; Highly similar to (defline not available 4929683) [H.		3.1	227	73	16.8
W01996	Hs.3945	H beta 58 homolog		3.1	359	118	2.5
AA393804	Hs.67052	ESTs; Highly similar to rap2 gene product [H.sapiens]		3.1	234	76	8.6
AA235289 D63477	Hs.247630	Human mRNA for KIAA0143 gene; partial cds		3.1	147	48	12.7
	Hs.84087	Human DNA sequence from PAC 196E23 on chromosom		3.1	31	1	2.3
AC000115	Hs.9030	•		3.1	31	1	2.6
U84011	Hs.904	amylo-1;6-glucosidase; 4-alpha-glucanotransferase (glyco		3.1	31	1	2.3
M74093	Hs.9700	cyclin E1		3.1	31	10	1.7
N75308	Hs.99433	ESTs	AAC1	3.1	94	30	5.8
X54925	Hs.83169	matrix metalloproteinase 1 (interstitial collagenase)	AACI	3.1	815	266	1.7
X12876	Hs.65114	keratin 18		3.1	68	23	2.8
M34309	Hs.199067	v-erb-b2 avian erythroblastic leukemia viral oncogene hom	BCF1	3	30	23 10	0.9
AA293300	Hs.9598	ESTs; Weakly similar to semaphorin C [M.musculus]	BCR2	3	816	275	3.9
AA609773	Hs.250175	Homo sapiens clone 23904 mRNA sequence		3	380	127	5.5
AA505133	Hs.62273	ESTs	CAA2	3	594	201	2.3
HG2981-HT3127	11- 70000	"Epican, Alt. Splice 11"	DOES	2.9	114	39	9.9
AA195936	Hs.76362	general transcription factor IIA; 1 (37kD and 19kD subunit				39 74	
AA419622	Hs.104800	ESTs	BCN1	2.9	214		3.7
R53457	Hs.26040	ESTs	BCX1	2.8	751	270	1.3
AA234561	Hs.22862	ESTs	BCZ1	2.8	131	47	3.9
AA428062	Hs.98558	ESTs	BCU7	2.7	864	321	0.6
AA620795	Hs.8207	ESTs	BCQ8	2.5	392	155	4.3
AA449749	Hs.31386	ESTs; Highly similar to secreted apoptosis related protein		2.1	1561	757	1.7
C13992	Hs.93668	ESTs	BCQ7	1.8	1047	596	1.6
H85169	Hs.172455	solute carrier family 5 (inositol transporters); member 3	BCW2	1	1	1	1

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	I IniCono ID			ratio tumor/	90%tile	75%tile	ratio tumor/
Accession	UniGene ID	UniGene Title		body		body	
AA126474	Hs.155223	stanniocalcin 2		72.2	722	1	1.9
AA434329	Hs.36563		3CJ7	40.2	402	1	4
AA250737	Hs.72472		3CY2	35.9	359	10	29.7
X82153	Hs.83942	cathepsin K (pycnodysostosis)		34.3	411	12	5.1
X03635	Hs.1657	* * * * * * * * * * * * * * * * * * * *	3CQ3	32.2	322	1	4.7
H09290	Hs.76550	ESTs; Weakly similar to unknown [H.sapiens]		30.6	306	4	26.5
AA428090	Hs.26102	ESTs B	3CN2	29	290	1	26.8
AA419547	Hs.11713	ESTs		26.3	356	14	1
AA256485	Hs.182471		3CO2	25.4	508	20	3
N67239	Hs.10760	ESTs B	3CX9	25.1	288	12	6.7
Z38595	Hs.125019	ESTs; Weakly similar to !!!! ALU SUBFAMILY J WARNING B	BCY3	24.2	242	10	5.6
H25836	Hs.83429	tumor necrosis factor (ligand) superfamily; member 10		22.8	228	9	12.4
HG1763-HT1780		Prolactin-Induced Protein		22.7	760	34	1.4
AA411621	Hs.8895	ESTs		21.2	212	6	17.4
N46252	Hs.29724	ESTs B	BCX6	20.9	209	1	19.5
U05237	Hs.99872	fetal Alzheimer antigen		20.6	206	4	19.1
U48807	Hs.2359	dual specificity phosphatase 4		20.2	202	5	1.3
AA070801	Hs.51615	ESTs; Weakly similar to !!!! ALU SUBFAMILY SQ WARNI		18.7	187	1	17
U28831	Hs.44566	Human protein immuno-reactive with anti-PTH polyclonal an	tibodie	s 18.6	186	10	1.5
AA292066	Hs.240802	ESTs; Weakly similar to Br140 [H.sapiens]		17.5	175	2	12.8
N26722	Hs.42645	ESTs B	BCY5	17.4	174	9	6.9
AA291725	Hs.105700		3CX2	17.4	409	24	7.8
AA065217	Hs.169674	ESTs		16.2	162	1	
D13666	Hs.136348		BCA4	15.7	1030	66	
AA621169	Hs.8687		3CX8	15.6	156	7	
L07615		"Human neuropeptide Y receptor Y1 (NPYY1) mRNA, exo		15.3	153	1	14.1
AA007234	Hs.30098	ESTs		14.9	149	1	6.4
F01831	Hs.14838	ESTs B	3CX4	14.6	219	15	7.6
N66818	Hs.42179	ESTs B	3CY6	14.5	145	1	2.4
H05509	Hs.24639	ESTs		14.2	142	1	9.5
AA149007	Hs.243954	ESTs		13.7	137	1	8.9
D12485	Hs.11951	phosphodiesterase I/nucleotide pyrophosphatase 1 (homo B	3CA2	13.2	244	19	9.9
AA490262	Hs.15485	ESTs; Moderately similar to APXL gene product [H.sapien B	BCU8	13.2	331	25	12.4
W93640	Hs.4779	ESTs		13.1	131	1	5.1
AA458761	Hs.18387	ESTs		12.7	311	25	2.4
Z48633	Hs.6940	H.sapiens mRNA for retrotransposon		12.4	124	6	10.8
AA227219	Hs.110826	Homo sapiens CAGF9 mRNA; partial cds		12.3	123	1	11.3
T97307	Hs.161720	ESTs; Moderately similar to !!!! ALU SUBFAMILY J WARN		12.3	129	11	11.7
D31352	Hs.31433	ESTs; Weakly similar to !!!! ALU SUBFAMILY SQ WARNI		11.7	117	1	10.1
AA251089	Hs.94576	ESTs; Weakly similar to phosducin; retinal [H.sapiens]		11.5	115	1	6.9
AA224180	Hs.187579	ESTs; Weakly similar to !!!! ALU SUBFAMILY J WARNING		11.5	115	1	
F11019	Hs.12696	ESTs		11.4	114	1	
X17059	Hs.155956	N-acetyltransferase 1 (arylamine N-acetyltransferase)		10.8	706	66	
H93575	Hs.227146	ESTs		10.5	105	1	
F13673	Hs.99769		3CN4	10.4	880	85	
AA411745	Hs.239681	ESTs; Weakly similar to KIAA0554 protein [H.sapiens]		10.3	103	1	
M90516	Hs.1674	glutamine-fructose-6-phosphate transaminase 1		10	100	1	
AA425309	Hs.33287		3CQ1	9.9	483	49	
AA487468	Hs.100686	ESTs; Moderately similar to secreted cement gland protei	3CX3	9.9	351	36	
M23379	Hs.758	RAS p21 protein activator (GTPase activating protein) 1		9.6	96	1	
T25867	Hs.7549		BCY9	9.6	124	13	
R63542	Hs.110488	ESTs		9.5	95	1	
M69225	Hs.620	bullous pemphigoid antigen 1 (230/240kD)		9.4	94	1	
Al039722	Hs.171205	ESTs		9.4	94	3	
U18321	Hs.159627	Death associated protein 3		9.3	93	5	
AA283006	Hs.50758	chromosome-associated polypeptide C		9.3	93	1	
U44378	Hs.75862	MAD (mothers against decapentaplegic; Drosophila) hom		9.3	93	1	7.8

AA235112	Hs.106227	ESTs; Moderately similar to similar to murine RNA-binding		9.1	91	1	7.6
AA028028	Hs.61460	ESTs	BCX5	9	90	1	5.5
M77142	Hs.239489	TIA1 cytotoxic granule-associated RNA-binding protein		8.9	89	5	8
HG2981-HT3125	11- 40450	"Epican, Alt. Splice 1"		8.5	85	1	3.2
U33147	Hs.46452	mammaglobin 1		8.5	2058	243	1.4
AA280036	Hs.145374	ESTs; Weakly similar to W01A6.c [C.elegans]	2011	8.5	127	15	1.6
AA609200	Hs.162686	ESTs	BCY4	8.5	85	1	4.3
D83004	Hs.75355	ubiquitin-conjugating enzyme E2N (homologous to yeast L	,	8.5	85	1	7.2
AA257971	Hs.21214	ESTs		8.3	83	3	1.8
U59863	Hs.146847	TRAF family member-associated NFKB activator		8.2	82	1	6.8
T16387	Hs.65328	ESTs		8.2	82	1	6.4
H13108	Hs.107968	ESTs ESTs		8.2	82	1	7.4
D31161	Hs.68613			8.1	81	1	4.6
M28213	Hs.78305	RAB2; member RAS oncogene family		7.8	78 77	1	5.6
AA031357	Hs.31803	ESTs		7.7	77	1	5.1
N66857	Hs.14808	ESTs; Weakly similar to !!!! ALU CLASS C WARNING EN		7.7	77	1	5
AA416997	Hs.59622	ESTs		7.6	144	19	13.9
R34531	Hs.243068	KIAA0480 gene product	D4.40	7.6	76	1	5
W37145	Hs.30029	ESTs	PAA9	7.5	136	18	3.4
D60237	Hs.14368	SH3-binding domain glutamic acid-rich protein like		7.5	75	1	6.5
AA054228	Hs.23165	ESTs		7.4	74	1	6
M99701	Hs.95243	transcription elongation factor A (SII)-like 1		7.3	73	1	5.3
Z46629	Hs.2316	SRY (sex-determining region Y)-box 9 (campomelic dyspl		7.3	73	1	5.2
AA478446	Hs.69559	ESTs; Weakly similar to Bat2 [H.sapiens]		7.2	72	1	5.7
U83908	Hs.247134	Homo sapiens nuclear antigen H731-like protein mRNA; c		7.2	72	1	5.8
AA199828	Hs.188662	ESTs	BCY8	7.1	71	1	6.5
D61676	Hs.21851	Homo sapiens mRNA; cDNA DKFZp586J2118 (from clone	BCB9	7.1	392	56	3.6
AA211941	Hs.109643	polyadenylate binding protein-interacting protein 1		7.1	71	1	6.2
H18027	Hs.184697	receptor for virally-encoded semaphorin	BCY7	7.1	150	21	14.5
AA032147	Hs.23296	ESTs		7	70	1	6.5
AA292701	Hs.5364	ESTs		6.9	69	1	4.4
J04088	Hs.156346	topoisomerase (DNA) II alpha (170kD)		6.8	68	1	5.6
Z38763	Hs.15740	ESTs		6.7	67	1	6.3
Al287461	Hs.164950	ESTs		6.7	67	1	6
AA195260	Hs.204151	ESTs; Moderately similar to !!!! ALU SUBFAMILY SX WAF	₹	6.7	67	1	5.7
N29888	Hs.169539	ESTs		6.5	65	4	5.3
AA490862	Hs.55901	ESTs; Moderately similar to !!!! ALU SUBFAMILY SQ WA		6.5	65	1	5.6
U37519	Hs.87539	aldehyde dehydrogenase 8		6.4	428	67	2.3
W23625	Hs.8739	ESTs		6.4	64	1	5.1
AA609723	Hs.30652	ESTs	PAA3	6.3	63	1	5.4
H10933	Hs.10067	ESTs	BCA1	6.3	693	110	7.2
N67711	Hs.151046	Homo sapiens clone 23859 mRNA sequence		6.3	63	1	5.8
AA430124	Hs.234607	ESTs		6.2	62	1	5.4
U35835	Hs.155637	protein kinase; DNA-activated; catalytic polypeptide		6.1	61	1	5.7
AA262491	Hs.186572	ESTs		5.8	58	1	5
D60302	Hs.108977	ESTs	BCY1	5.8	321	55	17
R51309	Hs.70823	KIAA1077 protein	BCN5	5.7	567	100	6.7
AA092376	Hs.90606	15 kDa selenoprotein		5.7	57	1	5
U90914	Hs.5057	carboxypeptidase D		5.6	56	1	5.3
AA130273	Hs.7584	ESTs; Weakly similar to (defline not available 4240269) [H		5.5	55	1	5.2
AA491465	Hs.28792	ESTs	BCU9	4.7	381	81	6.4
AA047036	Hs.62817	ESTs	BCR9	4.6	427	93	10.4
U41060	Hs.79136	Human breast cancer; estrogen regulated LIV-1 protein (L		4.5	1472	330	2.1
AA169379	Hs.72865	ESTs	BCU4	4.1	334	82	3.4
N22107	Hs.172241	ESTs; Moderately similar to !!!! ALU SUBFAMILY SC WAF		3.9	322	83	4.4
AA609651	Hs.112742	ESTs	BCX7	3.9	60	16	4.8
N91023	Hs.170057	ESTs	BCW4	3.8	425	111	4
C00038	Hs.23579	ESTs	BCQ5	3.8	585	153	3.7
N68921	Hs.34806	ESTs; Weakly similar to neogenin [H.sapiens]	BCW3	3.6	402	112	4.9
W72838	Hs.58213	ESTs	BCH1	3.5	2073	595	2.1

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AA609773	Hs.250175	Homo sapiens clone 23904 mRNA sequence	BCR2	3	816	275	3.9
AA419622	Hs.104800	ESTs	BCN1	2.9	214	74	3.7
AA234561	Hs.22862	ESTs	BCZ1	2.8	131	47	3.9
AA620795	Hs.8207	ESTs	BCQ8	2.5	392	155	4.3
C13992	Hs.93668	ESTs	BCQ7	1.8	1047	596	1.6
H85169	Hs.172455	solute carrier family 5 (inositol transporters); member 3	BCW2	1	1	1	1

(CONT.)

Accession	UniGene ID	UniGene Title		ratio tumor <i>i</i> body	90%tile turnor	75%tile body	ratio tumor/ normal breast
W72838	Hs.58213	ESTs	BCH1	3.5	2073	595	2.1
D12485	Hs.11951	phosphodiesterase I (PC-1)	BCA2	13.2	244	19	9.9
AA434329	Hs.36563	ESTs	BCJ7	40.2	402	1	4
AA419622	Hs.104800	ESTs	BCN1	2.9	214	74	3.7
R51309	Hs.70823	KIAA1077 protein	BCN5	5.7	567	100	6.7
AA256485	Hs.182471	ESTs	BCO2	25.4	508	20	3
C00038	Hs.23579	ESTs	BCQ5	3.8	585	153	3.7
	Hs.250175	Homo sapiens clone 23904 mRNA sequence	BCR2	3	816	275	3.9
AA609773		secreted frizzled-related protein 4	BCX2	17.4	409	24	7.8
AA291725 Z38595	Hs.105700 Hs.125019	ESTs	BCY3	24.2	242		5.6

GGACTTGCCCTAACAGAGCCTCAACAACTACCTGGTGATTCCTACTTCAGCCCCTTGGTGTGAGCAGCTTCTCAACATGA GAGGTCGGCAGAAGTGACAAGCTTTCCCTGCCTGGCTTTGAGAACCTCACAGCAGGATATAACAAATTTCTCAGGCCCAA TTTTGGTGGAGAACCCGTACAGATAGCGCTGACTCTGGACATTGCAAGTATCTCTAGCATTTCAGAGAGTAACATGGACT ACACAGCCACCATATACCTCCGACAGCGCTGGATGGACCAGCGGCTGGTGTTTGAAGGCAACAAGAGCTTCACTCTGGAT GGGAAACAGGCTCATCCGCCTCTTCTCCAATGGCACGGTCCTGTATGCCCTCAGAATCACGACAACTGTTGCATGTAACA TGGATCTGTCTAAATACCCCATGGACACACAGACATGCAAGTTGCAGCTGGAAAGCTGGGGCTATGATGGAAATGATGTG GAGTTCACCTGGCTGAGAGGGAACGACTCTGTGCGTGGACTGGAACACCTGCGGCTTGCTCAGTACACCATAGAGCGGTA TTTCACCTTAGTCACCAGATCGCAGCAGGAGACAGGAAATTACACTAGATTGGTCTTACAGTTTGAGCTTCGGAGGAATG GATTCAGTCCCTGCAAGAACCTGCATTGGAGTGACGACCGTGTTATCAATGACCACACTGATGATCGGGTCCCGCACTTC TCTTCCCAACACCAACTGCTTCATCAAGGCCATCGATGTGTACCTGGGGGATCTGCTTTAGCTTTGTGTTTGGGGCCCTTGC TAGAATATGCAGTTGCTCACTACAGTTCCTTACAGCAGATGGCAGCCAAAGATAGGGGGGACAACAAAGGAAGTAGAAGAA GTCAGTATTACTAATATCATCAACAGCTCCATCTCCAGCTTTAAACGGAAGATCAGCTTTGCCAGCATTGAAATTTCCAG CGACAACGTTGACTACAGTGACTTGACAATGAAAACCAGCGACAAGTTCAAGTTTGTCTTCCGAGAAAAGATGGGCAGGA TTGTTGATTATTTCACAATTCAAAACCCCAGTAATGTTGATCACTATTCCAAACTACTGTTTCCTTTGATTTTATGCTA GCCAATGTATTTTACTGGGCATACTACATGTATTTTTGAGTCAATGTTAAATTTCTTGCATGCCATAGGTCTTCAACAGG ACAAGATAATGATGTAAATGGTATTTTAGGCCAAGTGTGCACCCACATCCAATGGTGCTACAAGTGACTGAAATAATATT TGAGTCTTTCTGCTCAAAGAATGAAGCTCCAACCATTGTTCTAAGCTGTGTAGAAGTCCTAGCATTATAGGATCTTGTAA TAGAAACATCAGTCCATTCCTCTTTCATCTAATCAAGGACATTCCCATGGAGCCCAAGATTACAAATGTACTCAGGGCT GTTTATTCGGTGGCTCCCTGGTTTGCATTTACCTCATATAAAGAATGGGAAGGACCATTGGGTAACCCTCAAGTGTCA GAAGTTGTTTCTAAAGTAACTATACATGTTTTTTACTAAATCTCTGCAGTGCTTATAAAATACATTGTTGCCTATTTAGG GAGTAACATTTTCTAGTTTTTCTGGTTAAAATGAAATATGGGCTTATGTCAATTCATTGGAAGTCAATGCACTAAC TCAATACCAAGATGAGTTTTTAAATAATGAATATTATTTAATACCACAACAGAATTATCCCCAATTTCCAATAAGTCCTA TCATTGAAAATTCAAATATAAGTGAAGAAAAAATTAGTAGATCAACAATCTAAACAAATCCCTCGGTTCTAAGATACAAT GGATTCCCCATACTGGAAGGACTCTGAGGCTTTATTCCCCCACTATGCATATCTTATCATTTTATTATTATACACACATC AATATGGGCTGTTGCCATGAAGGCTTGCAGAATTGAGTCCATTTTCTAGCTGCCTTTATTCACATAGTGATGGGGTACTA AAAGTACTGGGTTGACTCAGAGAGTCGCTGTCATTCTGTCATTGCTGCTACTCTAACACTGAGCAACACTCTCCCAGTGG TGCTTGACCCAGGAACAAGTGGCTTAGCTTAAGTAAACTTGGCTTTGCTCAGATCCCTGATCCTTCCAGCTGGTCTGCTC TGAGTGGCTTATCCCGCATGAGCAGGAGCGTGCTGGCCCTGAGTACTGAACTTTCTGAGTAACAATGAGACACGTTACAG AGTACCAAAAGTGATTTTTGAGTGTGCCAGGGTAAAGGCTTCCAGTTCAGCCTCAGTTATTTTAGACAATCTCGCCATCT TTAATTTCTTAGCTTCCTGTTCTAATAAATGCACGGCTTTACCTTTCCTGTCAGAAATAAACCAAGGCTCTAAAAGATGA TTTCCCTTCTGTAACTCCCTAGAGCCACAGGTTCTCATTCCTTTTCCCATTATACTTCTCACAATTCAGTTTCTATGAGT TATGAGCCAATCATATTTGTGATTTTTTAAAAAAAGTTTAAAAGGAAATATCTGTTCTGAAACCCCACTTAAGCATTGTT 

#### FIGURE 13

 $\underline{\mathtt{ATG}}\mathtt{AACTACAGCCTCCACTTGGCCTTCGTGTGTCTGAGTCTCTTCACTGAGAGGATGTGCATCCAGGGGAGTCAGTTCAA$ CGTCGAGGTCGGCAGAAGTGACAAGCTTTCCCTGCCTGGCTTTGAGAACCTCACAGCAGGATATAACAAATTTCTCAGGC CCAATTTTGGTGGAGAACCCGTACAGATAGCGCTGACTCTGGACATTGCAAGTATCTCTAGCATTTCAGAGAGTAACATG CTGTGGGAAACAGGCTCATCCGCCTCTTCTCCAATGGCACGGTCCTGTATGCCCTCAGAATCACGACAACTGTTGCATGT AACATGGATCTGTCTAAATACCCCATGGACACAGACATGCAAGTTGCAGCTGGAAAGCTGGGGCTATGATGGAAATGA TGTGGAGTTCACCTGGCTGAGAGGGAACGACTCTGTGCGTGGACTGGAACACCTGCGGCTTGCTCAGTACACCATAGAGC GGTATTTCACCTTAGTCACCAGATCGCAGCAGGAGACAGGAAATTACACTAGATTGGTCTTACAGTTTGAGCTTCGGAGG AATGTTCTGTATTTCATTTTGGAAACCTACGTTCCTTCCACTTTCCTGGTGGTGTTGTCCTGGGTTTCATTTTGGATCTC TCTCGATTCAGTCCCTGCAAGAACCTGCATTGGAGTGACGACCGTGTTATCAATGACCACACTGATGATCGGGTCCCGCA CTTCTCTCCCAACACCAACTGCTTCATCAAGGCCATCGATGTGTACCTGGGGATCTGCTTTAGCTTTGTGTTTTGGGGCC TTGCTAGAATATGCAGTTGCTCACTACAGTTCCTTACAGCAGATGGCAGCCAAAGATAGGGGGACAACAAAGGAAGTAGA AGAAGTCAGTATTACTAATATCATCAACAGCTCCATCTCCAGCTTTAAACGGAAGATCAGCTTTGCCAGCATTGAAATTT CCAGCGACAACGTTGACTACAGTGACATGAAAACCAGCGACAAGTTCAAGTTTGTCTTCCGAGAAAAGATGGGC AGGATTGTTGATTATTTCACAATTCAAAACCCCAGTAATGTTGATCACTATTCCAAACTACTGTTTCCTTTGATTTTAT GCTAGCCAATGTATTTTACTGGGCATACTACATGTATTTT<u>TGA</u>

#### FIGURE 15

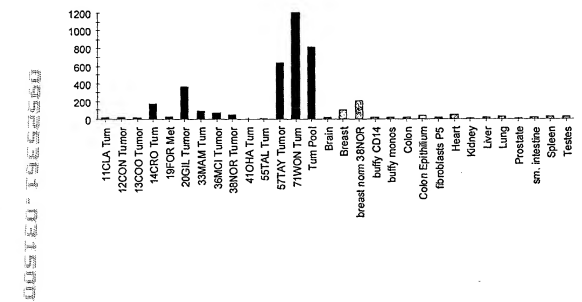
MNYSLHLAFVCLSLFTERMCIQGSQFNVEVGRSDKLSLPGFENLTAGYNKFLRPNFGGEPVQIALTLDIASISSISESNM DYTATIYLRQRWMDQRLVFEGNKSFTLDARLVEFLWVPDTYIVESKKSFLHEVTVGNRLIRLFSNGTVLYALRITTTVAC NMDLSKYPMDTQTCKLQLESWGYDGNDVEFTWLRGNDSVRGLEHLRLAQYTIERYFTLVTRSQQETGNYTRLVLQFELRR NVLYFILETYVPSTFLVVLSWVSFWISLDSVPARTCIGVTTVLSMTTLMIGSRTSLPNTNCFIKAID MAGGIEVE MED HYSSLQQMAAKDRGTTKEVEEVSITNIINSSISSFKRKISFASIEISSDNVDYSDLTMKTSDKFKFVFREKMG RIVDYFTIONPSNVDHYSKLLFPLIFMLANVFYWAYYMYF.

BCR3p1

Ac-Ala-Cys-Asn-Met-Asp-Leu-Ser-Lys-Tyr-Pro-Met-Asp-Thr-Gln-Thr-NH₂

Ac-Cys-Lys-Leu-Gln-Leu-Glu-Ser-Trp-Gly-Tyr -Asp-Gly-Asn-Asp-Vှal-NH₂

BCR3p2



breast cancer normal tissues

GTGAAGAGGCGCGCUTGACTGAGCTACGGTTCTGGCTGCGTCCTAGAGGCATCCGGGGCAGTAAAACCGCTGCGATC GCGGAGGCGGCGACGCCGAGACCAGGCCGGCAGGGGTGTCGGACGCAGGGCGCTGGGCCGGGTTTCGGCCTCGGCC ACAGCTTTTTTTCTCAAGGTGCAATGAAAGCCTTCCACACTTTCTGTGTTGTCCTTCTGGTGTTTTGGGAGTGTCTCTGAA GCCAAGTTTGATGATTTTGAGGATGAGGAGGACATAGTAGAGTATGATGATAATGACTTCGCTGAATTTGAGGATGTCAT GGAAGACTCTGTTACTGAATCTCCTCAACGGGTCATAATCACTGAAGATGATGAAGATGAGACCACTGTGGAGTTGGAAG GGCAGGATGAAAACCAAGAAGGAGATTTTGAAGATGCAGATACCCAGGAGGAGATACTGAGAGTGAACCATATGATGAT GAAGAATTTGAAGGTTATGAAGACAAACCAGATACTTCTTCTAGCAAAAATAAAGACCCAATAACGATTGTTGATGTTCC ACATCATTGGGAAGAATAAAAACAGTCGCCTTGCACAGGCCTGGTTTAACACTCATAGGGAGCCTTTTGGAGAGCAACTTT ACTTTAGTGGGGGATGATGGAACTAACAAAGAAGCCACAAGCACAGGAAAGTTGAACCAGGAGAATGAGCACATCTATAA CCTGTGGTGTTCTGGTCGAGTGTGCTGTGAGGGCATGCTTATCCAGCTGAGGTTCCTCAAGAGACAAGACTTACTGAATG ACCTAAGTCTGGAGCAAAGTATGGACTGCCGGACTCTTTGGCCATCCTGTCAGAGATGGGAGAAGTCACAGACGGAATGA TGGATACAAAGATGGTTCACTTCTTACACACCTATGCTGACAAGATTGAATCTGTTCATTTTTCAGACCAGTTCTCTGGT CCAAAAATTATGCAAGAGGAAGGTCAGCCTTTAAAGCTACCTGACACTAAGAGGACACTGTTGTTTACATTTAATGTGCC TGGCTCAGGTAACACTTACCCAAAGGATATGGAGGCACTGCTACCCCTGATGAACATGGTGATTTATTCTATTGATAAAG CCAAAAAGTTCCGACTCAACAGAGAGGCAAACAAAAAGCAGATAAGAACCGTGCCCGAGTAGAAGAACTTCTTGAAA TGAGGAAGATCCTGAGAAACAGCGCAGGCTGGAGGAGGCTGCATTGAGGCGTGACGAAAAAGAAGTTGGAAAAGAAGCAA ATGAAAATGAAACAAATCAAAGTGAAAGCCATGTAAAGCCATCCCAGAGATTTGAGTTCTGATGCCACCTGTAAGCTCTG **AATTCACAGGAAACATGAAAAACGCCAGTCCATTTCTCAACCTTAAATTTCAGACAGTCTTGGGCAACTGAGAAATCCTT** ATTTCATCATCTACTCTGTTTGGGGTTTGGGGTTTTACAGAGATTGAAGATACCTGGAAAGGGCTCTGTTTCAAGAATTT TTTTTTCCAGATAATCAAATTATTTTGATTATTTTATAAAAGGAATGATCTATGAAATCTGTGTAGGTTTTAAATATTTT AAAAATTATAATACAAATCATCAGTGCTTTTAGTACTTCAGTGTTTAAAGAAATACCATGAAATTTATAGGTAGATAACC TTGGAATTGCACAATAAACATTGCTTGATGTTTTCTTGTATGTCTACATTAAACTTGAGAAAAAGTAAAAATTAGAACAC TGTATGTAGTAATGAAATTTCAGGGACCCAGAACATAATGTAGTATATGTTTTTAGGTGGGAGATGCTGATAACAAAATT AATAGGAAGTCTGTAGGCATTAGGATACTGACATGTACATGGAAAATTCTAGGGACAGGAGCATCATTTTTTCCTTACCT GATACCACGAACCAGTGACAACGTGAATGCTGTATTTTAAGTGGTTGTATGTTTATTTTCTTGAGTAACAAATGCATGAA AAATTAATGCTTCACCTAGGTAAGATCATTGGTCTGTGTGAAATCACAAATGTTTTTTCCTTCTTGGTTGCAGCCTG GGTGGATGTTCATGGAGAAGCTCTGTTCTCTATATTATGGCTGTGCCGTTGCTTCTCCCTCTGCTTTTATCTTTTCCA CAGTTGAGGCTGGGTATGTTCTTTCAAAGAAATGGCCATGAATATGTGTAAGTATACTTTTGAAAATGAGCTTTCCTAAA CTATTGAGAGTTCTTTCCACCTCTTGCGGAACCAACTCTTGGAGGAGAGGCCCATGTATCTGCACGAGCACTTAGCTTGT TCAGATCTCTGCATTTTATAAATGCTTCTTACCAAGAAAGCATTTTTAGGTCATTGCTTGTACCAGGTAATTTTTGCCGG GGATGGGTAAGGGTTGGGTTTTCTGGTGGGAGTGGGGTGGTGGTATTTTTTTGTTGATGCTTTAGTGCAGGCCTGTTCTG AGGCAATAACAAGTTGCTGTGAAAACAGCATGTGCTGCTGCCTTTGTAACTGCATGGAAACTTTTCACATGGGTTTTTCT CCAAGTTAATACAGAAATATGTAAACTGAGAGATGCAAATGTAATATTTTTAACAGTTCATGAAGTTGTTATTAAAAATAA CTAACATAAAACTTAATTACTTTAATATTATAATTATAGTAGTGGCCTTGTTTTACAAACCTTTAAATTACATTTTAG AAATCAAAGTTGATAGTCTTAGTTATCTTTTGAGTAAGAAAAGCTTTCCTAAAGTCCCATACATTTGGACCATGGCAGCT AAAAAAAA

TGAGGAGGACATAGTAGAGTATGATGATAATGACTTCGCTGAATTTGAGGATGTCATGGAAGACTCTGTTACTGAATCTC CTCAACGGGTCATAATCACTGAAGATGATGAAGATGAGACCACTGTGGAGTTGGAAGGGCAGGATGAAAACCAAGAAGGA GATTTTGAAGATGCAGATACCCAGGAGGGAGATACTGAGAGTGAACCATATGATGATGAAGAATTTGAAGGTTATGAAGA CAAACCAGATACTTCTTCTAGCAAAAATAAAGACCCAATAACGATTGTTGATGTTCCTGCACACCTCCAGAACAGCTGGG AGTCGCCTTGCACAGGCCTGGTTTAACACTCATAGGGAGCTTTTGGAGAGCAACTTTACTTTAGTGGGGGATGATGGAAC TAACAAAGAAGCCACAAGCACAGGAAAGTTGAACCAGGAGAATGAGCACATCTATAACCTGTGGTGTTCTGGTCGAGTGT GCTGTGAGGGCATGCTTATCCAGCTGAGGTTCCTCAAGAGACAAGACTTACTGAATGTCCTGGCCCGGATGATGAGGCCA GTGAGTGATCAAGTGCAAATAAAAGTAACCATGAATGATGAAGACATGGATACCTACGTATTTGCTGTTGGCACACGGAA GACTGCCGGACTCTTTGGCCATCCTGTCAGAGATGGGAGAAGTCACAGACGGAATGATGGATACAAAGATGGTTCACTTC TTACACACCTATGCTGACAAGATTGAATCTGTTCATTTTTCAGACCAGTTCTCTGGTCCAAAAATTATGCAAGAGGAAGG TCAGCCTTTAAAGCTACCTGACACTAAGAGGACACTGTTGTTTACATTTAATGTGCCTGGCTCAGGTAACACTTACCCAA AGGATATGGAGGCACTGCTACCCCTGATGAACATGGTGATTTATTCTATTGATAAAGCCAAAAAGTTCCGACTCAACAGA GAAGGCAAACAAAAAGCAGATAAGAACCGTGCCCGAGTAGAAGAGAACTTCTTGAAACTGACACATGTGCAAAGACAGGA GCAGGCTGGAGGAGGCTGCATTGAGGCGTGACGAAAAAGAAGTTGGAAAAGAAGCAAATGAAAATGAAACAAATCAAAGT GAAAGCCATGTAAAGCCATCCCAGAGATTTGAGTTCTGA

MKAFHTFCVVLLVFGSVSEAKFDDFEDEEDIVEYDDNDFAEFEDVMEDSVTESPQRVIITEDDEDETTVELEGQDENQEG DFEDADTQEGDTESEPYDDEEFEGYEDKPDTSSKNKDPITIVDVPAHLQNSWESYYLETLMVTGLLAYIMNYIIGKNKN SRLAQAWFNTHRELLESNFTLVGDDGTNKEATSTGKLNQENEHIYNLWCSGRVCCEGMLIQLRFLKRQDLLNVLARMMRP VSDQVQIKVTMNDEDMDTYVFAVGTRKALVRLQKEMQDLSEFCSDKPKSGAKYGLPDSLAILSEMGEVTDGMMDTKMVHFLHTYADKIESVHFSDQFSGPKIMQEEGQPLKLPDTKRTLLFTFNVPGSGNTYPKDMEALLPLMNMVIYSIDKAKKFRLNR EGKQKADKNRARVEENFLKLTHVQRQEAAQSRREEKKRAEKERIMNEEDPEKQRRLEEAALRRDEKEVGKEANENETNQS ESHVKPSQRFEF.

# FIGURE 2 1

Peptide Name: BCQ8p1

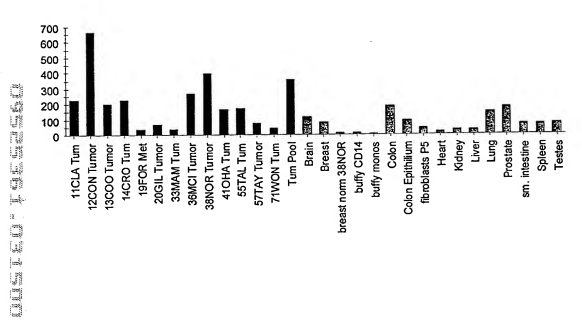
Sequence: Ac-Cys-Lys-Pro-Asp-Thr-Ser-Ser-Lys-Asn-Lys

-Asp-Pro-lle-Thr-NH₂

Peptide Name: BCQ8p2

Sequence: H-Lys-Phe-Asp-Asp-Phe-Glu-Asp-Glu-Glu-Asp

-Ile-Val-Glu-Tyr-Cys-NH₂



breast cancer normal tissues

GCGCCCGCGCTCGCAGGCCACTCTCTGCTGTCGCCCGTCCCGCGCGCTCCTCCGACCCGCTCCGCTCCGCTCCGCTCGGCCCCGCCCC CCCGTCAACATGATCCGCTGCGGCCTGGCCTGCGGCGCTGCCCCTGCTCCTACTCAGCGCCATCGCCTTCGACAT CATCGCGCTGGCCGGCGGGTTGCTGCAGTCTAGCGACCACGGCCAGACGTCCTCGCTGTGGTGGAAATGCTCCCAAGAGGGCGGCG GCAGCGGGTCCTACGAGGAGGGCTGTCAGAGCCTCATGGAGTACGCGTTGGGGTTAGAGCAGCGGCTGCCATGCTCTTCTGTGGCTTCATC ATCCTGGTGATCTGTTTCATCCTCCTTCTTCGCCCTCTGTGGACCCCAGATGCTTGTCTTCCTGAGAGTGATTGGAGGTCTCCTTGC CTTGGCTGCTGTTTCCAGATCATCTCCCTGGTAATTTACCCCGTGAAGTACACCCAGACCTTCACCCTTCATGCCAACCCTGCTGTCA CTTACATCTATAACTGGGCCTACGGCTTTGGGTGGCCACGACTTATCCTGATTGGCTGTCCTTCTTCTTCTGCTGCCTCCCCAAC GCTGAGATGGACTCCAGAAGAAGAAACTGTTTCTCCAGGCGACTTTGAACCCATTTTTTGGCAGTGTTCATATTATTAAACTAGTCAAA **AATGCTAAAATAATTTGGGAGAAAATATTTTTTAAGTAGTGTTATAGTTTCATGTTTATCTTTTATTATGTTTTGTGAAGTTGTGTCTT** TAACACTTTATAAGGTAAAAATGAGGTTTCCAAGATTTAATAATCTGAYCAAGTTCTTGTTATTTCCAAATAGAATGGACTCGGTCTGT GCCTTCGAACTATTTAAGGAAAGCAAAATCATTTCCTAAATGCATATCATTTGTGAGAATTTCTCATTAATATCCTGAATCATTC TAGCTAAGGCTTCATGTTGACTCGATATGTCATCTAGGAAAGTACTATTTCATGGTCCAAACCTGTTGCCATAGTTGGTAAGGCTTTCC tttaagtgtgaaatatttagatgaaattttctcttttaaagttctttatagggttagggtgtgggaaaatgctatattaataatctgt AGTGTTTTGTGTTTATATGTTCAGAACCAGAGTAGACTGGATTGAAAGATGGACTGGGTCTAATTTATCATGACTGATAGATCTGGTTA agttgtgtagtaaagcattaggagggtcattcttgtcacaaaagtgccactaaaacagcctcaggagaataaatgacttgctttctaa ATCTCAGGTTTATCTGGGCTCTATCATATAGACAGGCTTCTGATAGTTTGCAACTGTAAGCAGAAACCTACATATAGTTAAAATCCTGG TCTTTCTTGGTAAACAGATTTTAAATGTCTGATATAAAACATGCCACAGGAGAATTCGGGGATTTGAGTTTCTCTGAATAGCATATATA TGATGCATCGGATAGGTCATTATGATTTTTTACCATTTCGACTTACATAATGAAAACCAATTCATTTTAAATATCAGATTATTATTTTG 

## FIGURE 23

ATGATCCGCTGCGGCCTGGCCTGCAGCGCTGCCGCTGGATCCTGCCCCTGCTCCTACTCAGCGCCATCGCCTTCGACATCATCGCGCT
GGCCGGCCGGCGGCTGGTTGCAGTCTAGCGACCACGGCCAGACGTCCTCGCTGTGGTAAATGCTCCCAAGAGGGCGGCGCAGCGGGT
CCTACGAGGGGGGCTGTCAGAGCCTCATGGAGTACGCGTGGGGTAGAGCAGCGGCTGCCATGCTCTTCTTGTGGCTTCATCATCCTGGTG
ATCTGTTTCATCCTCTCCTTCTTCGCCCTTGTGGACCCCAGATGCTTGTCTTCCTGAGAGTGATTGGAGGTCTCCTTGCCTTGCCTTGCCTG
TGTTTCCAGATCATCTCCCTGGTAATTTACCCCGTGAAGTACACCCAGACCTTCACCTTCATCCAACCCTGCTGCACTTACATCT
ATAACTGGGCCTACGGCTTTGGGTGGGCAGCCACGATTATCCTGATTGGCTGTGCCTTCTTCTTCTGCTGCCTCCCCAACTACGAAGAT
GACCTTCTGGGCAATGCCAAGCCCAGGTACTTCTACACATCTGCCTTA

numan_BCQ5 nouse_BCQ5 cat_BCQ5	1 1 1	MIRCGLACERCRWILPLLLLSAIAFDIIALAGRGWLQSSDHGQTSSLWWK MLRCGLACERCRWILPLLLLSAIAFDIIALAGRGWLQSSNHIQTSSLWWR	50 50 0
numan_BCQ5	51	CSQEGGGSGSYEEGCQSLMEYAWGRAAAAMLFCGFIILVICFILSFFALC	100
nouse_BCQ5	51	CFDEGGGSGSYDDGCQSLMEYAWGRAAAATLFCGFIILCICFILSFFALC	100
cat_BCQ5	1	EYAWGRAAAATLFCGFIILVICFILSFFALC	31
numan_BCQ5	101	GPQMLVFLRVIGGLLALAAVFQIISLVIYPVKYTQTFTLHANPAVTYIYN	150
nouse_BCQ5		GPQMLVFLRVIGGLLALAAIFQIISLVIYPVKYTQTFRLHDNPAVNYIYN	150
rat_BCQ5		GPQMLVFLRVIGGLLALAAVFQIISLVIYPVKYTQTFRLHDNPAVNYIYN	81
numan_BCQ5 mouse_BCQ5 rat_BCQ5	151	WAYGFGWAATIILIGCAFFFCCLPNYEDDLLGNAKPRYFYTSA 193 WAYGFGWAATIILIGCSFFFCCLPNYEDDLLGAAKPRYFYPPA 193 WAYGFGWAATIILIGCSFFFCCLPNYEDDLLGNAKPRYFYTSA 124	

#### BCQ5p1

Ac-Cys-Ser-Tyr-Ser-Ala-Pro-Ser-Pro-Ser-Thr -Ser-Ser-Arg-Trp-Pro-NH₂

BCQ5p2

Ac-Cys-Leu-Pro-Asn-Tyr-Glu-Asp-Asp-Leu-Leu-Gly-Asn-Ala-Lys-Pro-NH₂

BCQ5p3

Ac-Cys-Gly-Gly-Asn-Ala-Pro-Lys-Arg-Gly-Gly-Gly-Gly-Arg-Gly-Ser-Tyr-NH₂

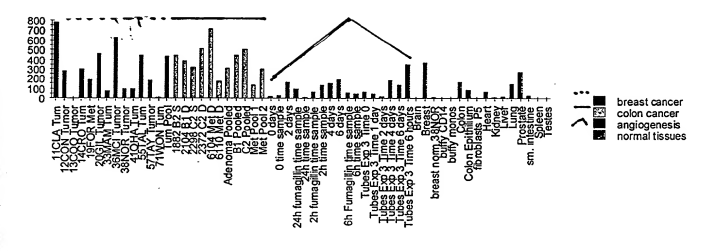


FIGURE 27

CGCTGCGGCCTGCCGAGCCGCTGCAGGTGGATCCTGCCCCTGCTGCTCAGCGCCATCGCCTTCGACATCATCGCGCTGGCCGGC CGCGGCTGGCTGCAGTCTAGCAACCACATCCAGACATCGTCGCTTTGGTGGAGGTGTTTCGACGAGGGCGGCGGCAGCGGCTCCTACGAC GATGGCTGCCAGAGCCTCATGGAGTACGCATGGGGACGAGCAGCTGCAGCCACGCTTTTCTGTGGCTTTATCATCCTGTGCATCTGCTTC ATTCTCTCGTTCTTCGCCCTGTGTGGACCCCAGATGCTTGTTTTCCTGAGAGTCATTGGAGGCCTCCTCGCACTGGCTGCCATATTCCAG ATCATCTCCCTGGTAATCTACCCCGTGAAGTACACACAGACCTTCAGGCTTCACGATAACCCTGCTGTTAATTACATCTATAACTGGGCC TATGGCTTCGGATGGGCGCCACCATCATCTTGATTGGTTGTTCCTTCTTCTTCTGCTGCCTCCCAACTACGAGGATGACCTTTTGGGG GCCGCCAAGCCCAGGTACHTCTATCCCCCAGCCTAATGTGGGAGGAAGAGCCTGAGAAAAGCCTGCTGCAAGATGGATCTGAGGAGGAAA CTGTTCTCCAAGGCACAAGGAACCTACGTTTGGGCAATGTTCATATGATCAGAAATGTTAGAATAAATGCTAAAGAAAATTCTTCATAAT TAGTGTTAAGTTTCATGTATGTCGTGTGGAGTTAAAAAGACTTGAATTCTGTTTGCTAAGTATATGCTAATTTTTCCTTATGTCAATTCT ATACCATTTAAGCTTCATTTGTTAAAGAATATGCCTGTGAAACTTGATAAGGTAGAAATGCAGCAGCCTCTCATTTAATAATCTGATGGG GAAATTAGTTTGTTTTTTTTTAAAAGACCTTATTTTGAGTTTTCAGTTACATAAAGAAGCAGAAGCAGATTGGTTTCCTAAGTGAGCATC ATTTGTGAGAATTTTTAGTCAGTGTTTTGAACAATTATTGTTTTTCTAAGCTTCATGTTGACTTTCTCTGATGCGTAGAAAAGTGTTCTA TGGTGGATGTGAAGAGGTCAGGTAGGAAGGCACAGGAGGGTCACCACTGTCACAGCAGTGCCATGCAGACATCCTAGGAGAAGACATGGC TTAAAAACCTGGTCTTCCTTGGTAAGCAGACTTAAAATATCTGTATAGTACATGCAAGTGGAAAATTTGGGAATGCGTGTCTCTGAATAC ATACCGGAAGGGCTACTATTACCTTTTCTTACCATTTATACTTACCTAATGGAAACGAGCTTGTTTTAACTATCAGAACACTATTTTGT 

## FIGURE 28

## FIGURE 30

GAATACGCCTGGGGCCGAGCAGCTGCTGCCACTCTTTTTGTGGATTCATCATCTGGTCATCTGCTTCATCCTCTCTTCTTCGCCCTG
TGTGGACCCCAGATGCTTGTTTTCCTGAGAGTGATTGGAGGCCTTCTCGCACTGGTTGTTTCCAGATCATCTCCCTGGTTATCTAT
CCCGTGAAGTACACAAAACCTTCAGGCTTCATGATAATCCCGCTGTTAATTACATCTACAACTGGGCCTATGGCTTCGGATGGCAGCC
ACGATCATCTTGATTGGTTGCTCTTTCTTCTTCTTCTGCTGCCTCCCCAACTACGAGGATGACCTTCTGGGCAATGCAAAGCCCAGGTACTTC
TATACATCTGCCTAA

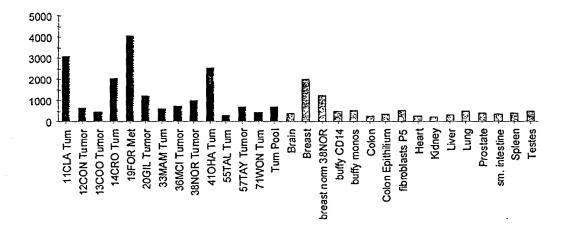
## FIGURE 33

## FIGURE 34

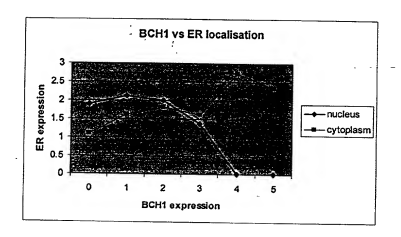
 ${\tt MKFLAVLVLLGVSIFLVSAQNPTTAAPADTYPATGPADDEAPDAETTAAATTATTAAPTTATTAASTTARKDIPVLPKWVGDLPNGRVCP.}$ 

H-Gln-Asn-Pro-Thr-Thr-Ala-Ala-Pro-Ala-Asp-Thr-Tyr-Pro-Ala-Cys-NH₂

Ac-Leu-Pro-Lys-Trp-Val-Gly-Asp-Leu-Pro-Asn-Gly-Arg-Val-Cys-Pro-OH

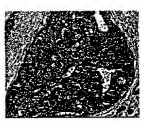


breast cancer normal tissues



## FIGURE38A

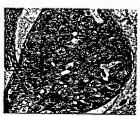
# FIGURE 38B





# FIGURE39A

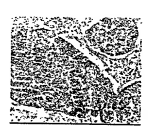
## FIGURE39B





# FIGURE 40A

# FIGURE 10B





## FIGURE 41 A

FIGURE 41B





**ÉGAGTTTTCTGGAGCTGTTGCAATGTGTATGCTGGTGAAATCTACTTGAGCATTAAGCAGTATCTCCCAGC ATTGTTAGCTACTGAGTGGCACATCTTCAGTACGCATGATTCGTGGGGGACTCAGGCAGAGGTAAAAGTGT ĞAAACTTTTCAGCATTACCTAAGAAGCAAAGGCTCAATTTTGGCTGCTTCATTCTTATCTCTTCTGCCACA** GTTCTAACGTGCCTGATCTACTGAGACCAAGG ACCAATGACTCAGAAGGGAAAATGGGATTTAAACAC <u>ECAAAGATCATGGGGAATTTCAGAGGTCATGCCCTCCCTGGAACCTTCTTTTTTATTATTGGTCTTTGGTG</u> GTGTACAAAGAGTATTCTGAAGTATATCTGCAAAAAGCAAAAGCGAACCTGCTATCTTGGTTCCAAAACAT TATTCTATCGATTGGAAATTTTGGAGGGAATTACAATAGTTGGCATGGCTTTAACTGGCATGGCTGGGGAG CAGTTTATTCCTGGAGGGCCCCATCTGATGTTATATGACTATAAACAAGGTCACTGGAATCAACTCCTGGG CTGGCATCATTTCACCATGTATTTCTTCTTTGGGCTGTTGGGTGTGGCAGATATCTTATGTTTCACCATCA GTTCACTTCCTGTGTCCTTAACCAAGTTAATGTTGTCAAATGCCTTATTTGTGGAGGCCTTTATCTTCTAC AACCACACTCATGGCCGGGAAATGCTGGACATCTTTGTGCACCAGCTGCTGGTTTTGGTCGTCTTTCTGAC AGGCCTCGTTGCCTTCCTAGAGTTCCTTGTTCGGAACAATGTACTTCTGGAGCTATTGCGGTCAAGTCTCA TTCTGCTTCAGGGGAGCTGGTTCTTTCAGATTGGATTTGTCCTGTATCCCCCCAGTGGAGGTCCTGCATGG GATCTGATGGATCATGAAAATATTTTGTTTCTCACCATATGCTTTTGTTGGCATTATGCAGTAACCATTGT AAGTTGAAAAAGACTTCTGTGCTGAACGAGAACAAGAATCAGAAGAAGAAATG CAGCTGGCTAAGGATGACTCTAAGTGTACTGTTTGCATTTCCAATTTGGTTAAAGTATTTGAATTTAAATA TTTTCTTTTTAGCTTTGAAAATATTTTGGGTGATACTTTCATTTTGCACATCATGCACATCATGGTATTCA TTTGCCTTATAGATATGCTCAAGGTTACTGGGCTTGCTACTATTTGTAACTCCTTGACCATGGAATTATAC 

### FIGURE 43

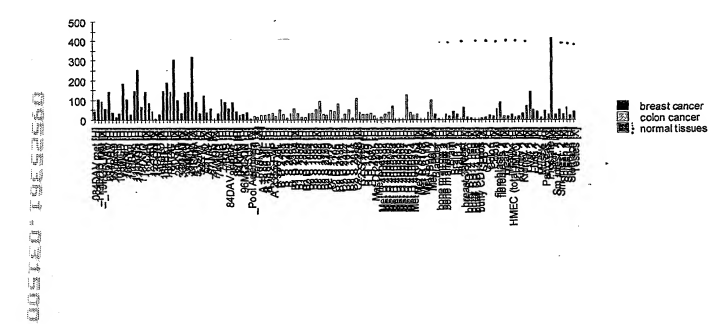
MTNDSEGKMGFKHPKIMGNFRGHALPGTFFFIIGLWWCTKSILKYICKKQKRTCYLGSKTLFYRLEILEGI TIVGMALTGMAGEQFIPGGPHLMLYDYKQGHWNQLLGWHHFTMYFFFGLLGVADILCFTISSLPVSLTKLM LSNALFVEAFIFYNHTHGREMLDIFVHQLLVLVVFLTGLVAFLEFLVRNNVLLELLRSSLILLQGSWFFQI GFVLYPPSGGPAWDLMDHENILFLTICFCWHYAVTIVIVGMNYAFITWLVKSRLKRLCSSEVGLLKNAERE QESEEEM.

#### BCN1p1

Ac-Tyr-Pro-Pro-Ser-Gly-Gly-Pro-Ala-Trp-Asp-Leu-Met-Asp-His-Cys-NH₂

BCN1p2

Ac-Cys-Leu-Lys-Asn-Ala-Glu-Arg-Glu-Glu-Glu-Ser-Glu-Glu-Glu-Met-OH



## FIGURE 46A

TTCCTCCGCGAAGGCTCCTTTGATATTAATAGTGTTGGTGTCTTGAAACTGACGTAATGCGCGG AGACTGAGGTCCTGACAAGCGATAACATTTCTGATAAAGACCCGATCTTACTGCAATCTCTAG  ${\tt CGTCCTCTTTTTGGTGCTGCTGGTTTCTCCAGACCTCGCGTCCTCTCGATTGCTCTCTCGCCTT}$ CCTATTTCTTTTTTTTTTTAAACAAAAACAACACCCCCTCCCCTCTCCCACCCGGCACCG CGGGCTGCCGAGGGTGTTCTTGACGATTAATCAACAGATGTACAGATCAGCTCTCAAAATGTC TTCTGTGTCTTCTGAGCGTCTTCTAAGACAATTGCATTAGCCTCCTGCTAGTTGACTAATAGAA TTAATAATTGTAAAAAGCACTCTAAAGCCACATGCCTTATGAAGTCAATGCTGGGTATGATTT TACAAAT GTCCGGAAAAAGAACCCCCCTCTGAGAAACGTTGCAAGTGAAGGCGAGGGCC AGATCCTGGAGCCTATAGGTACAGAAAGCAAGGTATCTGGAAAGAACAAAGAATTTTCTGCA GATCAGATGTCAGAAAATACGGATCAGAGTGATGCTGCAGAACTAAATCATAAGGAGGAACA TAGCTTGCATGTTCAAGATCCATCTTCTAGCAGTAAGAAGGACTTGAAAAGCGCAGTTCTGAG TGAGAAGGCTGGCTTCAATTATGAAAGCCCCAGTAAGGGAGGAAACTTTCCCTCCTTTCCGCA TGATGAGGTGACAGACAGAAATATGTTGGCTTTCTCATCTCCAGCTGCTGGGGGAGTCTGTGA GCCCTTGAAGTCTCCGCAAAGAGCAGAGGCAGATGACCCTCAAGATATGGCCTGCACCCCCTC AGGGGACTCACTGGAGACAAAGGAAGATCAGAAGATGTCACCAAAGGCTACAGAGGAAACA GGGCAAGCACAGAGTGGTCAAGCCAATTGTCAAGGTTTGAGCCCAGTTTCAGTGGCCTCAAA AAACCCACAAGTGCCTTCAGATGGGGGTGTAAGACTGAATAAATCCAAAACTGACTTACTGG TGAATGACAACCCAGACCCGGCACCTCTGTCTCCAGAGCTTCAGGACTTTAAATGCAATATCT GTGGATATGGTTACTACGGCAACGACCCCACAGATCTGATTAAGCACTTCCGAAAGTATCACT TAGGACTGCATAACCGCACCAGGCAAGATGCTGAGCTGGACAGCAAAATCTTGGCCCTTCAT AACATGGTGCAGTTCAGCCATTCCAAAGACTTCCAGAAGGTCAACCGTTCTGTGTTTTCTGGT GTGCTGCAGGACATCAATTCTTCAAGGCCTGTTTTACTAAATGGGACCTATGATGTGCAGGTG ACTTCAGGTGGAACATTCATTGGCATTGGACGGAAAACACCAGATTGCCAAGGGAACACCAA GTATTTCCGCTGTAAATTCTGCAATTTCACTTATATGGGCAACTCATCCACCGAATTAGAACAA CATTTTCTTCAGACTCACCCAAACAAAATAAAAGCTTCTCTCCCCTCTCTGAGGTTGCAAAAC CTTCAGAGAAAACTCTAACAAGTCCATCCCTGCACTTCAATCCAGTGATTCTGGAGACTTGG GAAAATGGCAGGACAAGATAACAGTCAAAGCAGGAGATGACACTCCTGTTGGGTACTCAGTG CCCATAAAGCCCCTCGATTCCTCTAGACAAAATGGTACAGAGGCCACCAGTTACTACTGGTGT AAATTTTGTAGTTTCAGCTGTGAGTCATCTAGCTCACTTAAACTGCTAGAACATTATGGCAAG CAGCACGGAGCAGTGCAGTCAGGCGGCCTTAATCCAGAGTTAAATGATAAGCTTTCCAGGGG CTCTGTCATTAATCAGAATGATCTAGCCAAAAGTTCAGAAGGAGAGACAATGACCAAGACAG ACAAGAGCTCGAGTGGGGCTAAAAAGAAGGACTTCTCCAGCAAGGGAGCCGAGGATAATATG GTAACGAGCTATAATTGTCAGTTCTGTGACTTCCGATATTCCAAAAGCCATGGCCCTGATGTA ATTGTAGTGGGGCCACTTCTCCGTCATTATCAACAGCTCCATAACATTCACAAGTGTACCATTA AACACTGTCCATTCTGTCCCAGAGGACTTTGCAGCCCAGAAAAGCACCTTGGAGAAATTACTT ATCCGTTTGCTTGTAGAAAAAGTAATTGTTCCCACTGTGCACTCTTGCTTCTGCACTTGTCTCC CGTAGATGTACTCCTCTTTCACTATGAAAGTGTGCATGAGTCCCAAGCATCGGATGTCAAACA AGAAGCAAATCACCTGCAAGGATCGGATGGGCAGCAGTCTGTCAAGGAAAGCAAAGAACACT CATGTACCAAATGTGATTTTATTACCCAAGTGGAAGAAGAGATTTCCCGACACTACAGGAGAG CACACAGCTGCTACAAATGCCGTCAGTGCAGTTTTTACAGCTGCCGATACTCAGTCACTACTGG AGCACTTCAACACTGTTCACTGCCAGGAACAGGACATCACTACAGCCAACGGCGAAGAGGAC GGTCATGCCATATCCACCATCAAAGAGGAGCCCAAAATTGACTTCAGGGTCTACAATCTGCTA AGAGAAGGACGGCTCAAAGAGAAAGTTTGGACCGAGAGTTCCAGTGATGACCTTCGCAATG CTGCTGACGCCTGTGTCTGGCACCCAAGAGCAGACAAAGACTCTAAGGGATAGTCCCAATGT GGAGGCCGCCCATCTGGCGCGACCTATTTATGGCTTGGCTGTGGAAACCAAGGGATTCCTGCA

## FIGURE 46B

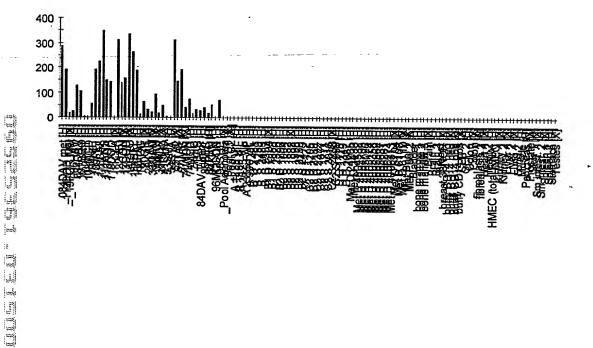
GGGGCGCCAGCTGGCGGAGAGAGTCTGGGGCCCTCCCCCAGCAGTATCCTGCATCGGGAG AAAACAAGTCCAAGGATGAATCCCAGTCCTGTTACGGAGGCGTAGAGGCTCCGGTGTTTTTT GTGCCAATTGCCTGACCACAAAGACCTCTCTCTGGCGAAAGAATGCAAATGGCGGATATGTAT GCAACGCGTGTGGCCTCTACCAGAAGCTTCACTCGACTCCCAGGCCTTTAAACATCATTAAAC AAAACAACGGTGAGCAGATTATTAGGAGGAGAACAAGAAAGCGCCTTAACCCAGAGGCACTT CAGGCTGAGCAGCTCAACAAACAGCAGAGGGGCAGCAATGAGGAGCAAGTCAATGGAAGCC CGTTAGAGAGGGGTCAGAAGATCATCTAACTGAAAGTCACCAGAGAGAAATTCCACTCCCC AGCCTAAGTAAATACGAAGCCCAGGGTTCATTGACTAAAAGCCATTCTGCTCAGCAGCCAGTC CTGGTCAGCCAAACTTTGGATATTCACAAAAGGATGCAACCTTTGCACATTCAGATAAAAAGT CCTCAGGAAAGTACTGGAGATCCAGGAAATAGTTCATCCGTATCTGAAGGGAAAGGAAGTTC TGAGAGAGGCAGTCCTATAGAAAAGTACATGAGACCTGCGAAACACCCAAATTATTCACCAC CAGGCAGCCCTATTGAAAAGTACCAGTACCCACTTTTTGGACTTCCCTTTGTACATAATGACTT CCAGAGTGAAGCTGATTGGCTGCGGTTCTGGAGTAAATATAAGCTCTCCGTTCCTGGGAATCC GCACTACTTGAGTCACGTGCCTGGCCTACCAAATCCTTGCCAAAACTATGTGCCTTATCCCACC TTCAATCTGCCTCCTCATTTTTCAGCTGTTGGATCAGACAATGACATTCCTCTAGATTTGGCGA TCAAGCATTCCAGACCTGGGCCAACTGCAAACGGTGCCTCCAAGGAGAAAACGAAGGCACCA CCAAATGTAAAAATGAAGGTCCCTTGAATGTAGTAAAAACAGAGAAAGTTGATAGAAGTAC TCAAGATGAACTTTCAACAAAATGTGTGCACTGTGGCATTGTCTTTCTGGATGAAGTGATGTA TGCTTTGCATATGAGTTGCCATGGTGACAGTGGACCTTTCCAGTGCAGCATATGCCAGCATCTT TGCACGGACAAATATGACTTCACAACACATATCCAGAGGGCCTGCATAGGAACAATGCACA AGTGGAAAAAATGGAAAACCTAAAGAG ATAGGTTTTCTTGATGGGAATTCAATAGCTTGTAATGTCTTATGAAGACCTATTAAAAAAATA AATCACTTTTTAAAACCTGTTAAGTCAAAACCATCTTGGCTAATATGTACTGGGGAAATAATC GAATTTGAAAGGGTTTACATATTATTATACTAAAGCAGTACTGGACTGGCCATTGGACCATTT GTTCCAAAACCCATAAATTGTTGCCTAAATTTATAATGATCATGAAACCCTAGGCAGAGGAGG AGAAATTGAAGGTCCAGGGCAATGAAAGAAAAATGGCGCCCTCTCAATTTAGTCTTCTCAT TGGCCATGTTTCAGATTTTGACCTAGAAATGCGAGCTGTGGTTAGGCTTGGTTAGAGTGCAGC GATTCATTTTGTCCACACATGTACCCATTTTAAAAAAACAATGTCCTCGATGCTTCTGTAGTGAT TTCATTTTAGCCAGGTATTTCTTTCTTGTGTGTGATGAACCAGTATGGATTTGCTTTTCTAAGCC TCCTGTTGGTTACTAATCTCACTTGGCACATTATAACTAAAGGAATCCCCTCAATTCAAAAGC ATAGATGGATACAAATGTCAGACCGTGGGTTTAATTTGTTTAGAACACATGGCATTTCTTCAC AAGGTAACCTGCTGTATTTATTTATTTTCTTTTGGTTAAATATAATTTCCAAACTTTGTGGTCAG GCAGCGTCTAAGGTTACCGTTACCACAGACTGACAGTTGGTATATGTACCAGCCAATCCCTTCA TTAAATGTATACAGATTTAGTTAAGTAGCATTAAATAGGATTCTTAGAAGTATGTCCTCATAG AACTTTTAATACTTAAGGCTTTGTAAAAACTATCCATGAAGGGAAAGCTCCTCAGCATAACTG CTCAGGGAAATAGGGCTAAATAACTGAACATTAAATAATTGGTTAAAGGTGCTGTTAGTCGA CCCAACCAGTAGTTTATTTTTTGCCACGGAGATGTAGAAGATATTACAAGCTACTGGATGCAC TTCTAGTAAATATTAATGTATTACATTTCAAATAATGGTGCCTGACATATTGAATAATTATTTT CTACAGTGTACGTATGCAACAAAGATATTCCATCATGCATTAGAGTCAGTTCTGGCTCTGCCT AGCTGTTTACATTTGCAAATGTAGCAAACAAGGTAATGAAGCAACTATTTCTATTGCAGTAGA TATCCTTTTGTGTGTGTGTGTGCATTAAAGTTGTAAACGGTAACATGAAACAAATGAAAGT TCTTGCTATAATGGTATGGAAAACAAGAAGGAAATGAAAATATTTTTATGCCTACTTAGGAAA TTGTTATGCTTAAGATGATAAACATATATCCTCTTTTTATTGCTTTGTCTATGTTTCATATGAAA GTTTTGTTTCTTTTCACAGTGCCGGGTCTTCGTTTCTTAAAGTTGGATGGCAGGTAGAGTTCA ACCAGTTCGTGACTGTTGTAGCGAATGAAGTTAAAAAAATGTCTTTCTGATGTTGTTGTCAT TTTCATTTTTGCATTTTTTGTTTGCATATTAAAAAAAAGAGAAAAGAGAAAGCAAGAGACAGA

# FIGURE 46C

AATCAGGACTAAGTCCTCTGCTTCAGTTTCATTGTTAACGGGCCTTATTCTGATCTCACCTGTC GCGTAGCTCTAATATTCACATAAACTGAAATAAAGAAGTGGAATGAGGAGCTTTGACATTCA GACTTGAGAGGTGACAATTAAAGATCTAAAAAAGAGAGGAGATTCCCCCAAACAACAATATT TAATTTTCTTAGTAAAAAGAATAACAGAATGCATCGTGGCAATCCTTAAGCAACATTATCTAT GTGGACTGCTTAAATCAGCAAAACACCAGAAGTTTGGTTAACTTGGGCAATATGACAAGTATT ACTTTTTGGGCAAAACTACTCATTAAGCAATTTCTCTAGTGTGTCGGACACAAATAGGTTCTTT TATCAACTAGCATTGGAAAATACATATCACTATTCTTGGAATATTTATGGTCAGTCTACTTTTT AGTAAAATATTTTTGGATAGCGTTGACACGATAGATCTTATTCCATACTTCTTTATTATTGATA ATTTTATTTTCATTTTTGCTTTCATTATTATACATATTTTGGTGGAGAAGAGGTTGGGCTTTTT TGAAAGAGACAAAATTTATTATAACACTAAACACTCCTTTTTTGACATATTAAAGCCTTTATT CCATCTCTCAAGATATATATAAAATTTATTTTTTTAATTTAAGATTTCTGAATTATTTTATCTT AAATTGTGATTTTAAACGAGCTATTATGGTACGGAACTTTTTTTAATGAGGAATTTCATGATGA TTTAGGAATTTTCTCTCTTGGAAAAGGCTTCCCCTGTGATGAAAATGATGTGCCAGCTAAAATT GTGTGCCATTTAAAAACTGAAAATATTTTAAAATTATTTGTCTATATTCTAAATTGAGCTTTGG ATCAAACTTTAGGCCAGGACCAGCTCATGCGTTCTCATTCTTCCTTTTCTCACTCTTTCTCACT CACTCACCTCTGTATTCATTCTGTTGTTTGGGATAGAAAAATCATAAAGAGCCAACCCATCTC AGAACGTTGTGGATTGAGAGAGACACTACATGACTCCAAGTATATGAGAAAAGGACAGAGCT CTAATTGATAACTCTGTAGTTCAAAAGGAAAAGAGTATGCCCAATTCTCTCTACATGACATAT TGAGATTTTTTTAATCAACTTTTAAGATAGTGATGTTCTGTTCTAAACTGTTCTGTTTTAGTGA AGGTAGATTTTTATAAAACAAGCATGGGGATTCTTTTCTAAGGTAATATTAATGAGAAGGGAA AAAAGTATCTTTAACAGCTCTTTGTTGAAGCCTGTGGTAGCACATTATGTTTATAATTGCACAT TTTAAAATGCCTGAGGAAATACATTTTTCTTAATAAACTGAAGAGTCTCAGTATGGCTATTAA AATAATTATTAGCCTCCTGTTGTGTGGCTGCAAAACATCACAAAGTGACCGGTCTTGAGACCT GTGAACTGCTGCCCTGTTTAGTAAATAAAATTAATGCATTTCTAGAGGGGGAATATCTGCCAT CCAGTGGTGGAAATGTGGAGTAAAGAAGCTGGTGGTCTGCTTCTGTGCTGTATGCCAGCCTTT TGCCTTAAGTTGAGAGGAGGTCAACTTTAGCTACTGTCTTTGGTTTGAGAGCCATGGCAAAAA AAAAAAAAGAAAAAAGATCAAGTCGTCTTTGGTGAGCCAGTAAGGTGAAAGCTTGCTGACT GTCCAAGGCACAAGAGAAAATTGAGGAATTGAAATGCAACCTGAGTATCAAACTAAATATTC TAATCAAAGGTAGGTACTGTTAGGTGGAATTCTATCAGCAGGCAACTGCAAATGAGAAGAAG GGGCAGAGGCATGGATTCTTTGCAGAGCACTTCCTTTTGGTTTTTCAGTACTGTTTCATAGACA GTGGGCTCACATGTTCCTGATAGTGCTGCAGTTGCTTAGAAAGCATCCCAGTTAATTĞCAGTA ATTAGAACTTCTGGAATATGCTAGGGCAGAAGTATGTCAAGTATGTCACATGAAGAAAATGT GAAATTCAAGAGTAATCCACACGTGAGAAACTAGACAATGTACATTCATGTTCTCTTGAAA GGAAAGGGAGAGCTGTAAGCTTCACTCTGTCCTACACCGGAGAAAAGCAGGAATAACTTTAC CGTGGAAATAATGTTTAGCTTTTATCAGAGAAAATTGTCCTTCTAGAGCATAGAGTCCCAAAA ATCACTTTTTCTTTTTGTGCCTCAGGTTCCTCASCTGTAAAATTGAAAAATATATGTATTA CTTGGTTGAAAGGCACCATAGGAGTGCCAAGTATTATTATGTGGCCAAGGGGGTTATTT AAACTGTCAGTTCCCAAAGGCCAGGAAAGGTTGGGGTCATTTTCTTAAAGACGAGCTG TAAATATCAACTAGGCAGCCAATAGTGTTGACTATGAAGATGCAAAACTATTACTAGGCT GATAAAATCATAGTTTCTTAATGGCTACCAATAAGGCAAATATCACAATAATAAACGCCA **AATTCCTTAGGGCGGACTATTTGACAACCACATGGAAAACTTTGGGGGAGGCATGAGGG** GGGAACATCTCAAAATGCCAATGTAAAATTTAACTTACAGCAATATTCACCAGCAGAAAA TGTCTTTCATATGGAATGATTTCATGTTGCTAAGAAAAGAATTCAATTTGTAGTCCTGA TTTGAATACTAGAATGTTGGCTATAATAGTTCTGTTCTTACAACACATGAAATTTTTTCGT TTTATTTTATTTTGTTTTCATAGTGCATGTTCATTTCTACTCACAAACATGTTCTTGGTGT **ATTTCTTATGCAAACAATCTTCAGGCAGCAAAGATGTCTGTTACATCTAAACTTGAATAA** TAAAGTTTTACCACCAGTTACACATAACGGCGTTGGTATGGTTTATATGGATTCACTTTC TCTGCTGAANTGAAATGGTGAAATCAAATCACCATTCTAAAAAAATTATTACTTATATTGA 

# FIGURE 46D

MVRKKNPPLRNVASEGEGQILEPIGTESKVSGKNKEFSADQMSENTDQSDAAELNHKEEHSLHVQ DPSSSSKKDLKSAVLSEKAGFNYESPSKGGNFPSFPHDEVTDRNMLAFSSPAAGGVCEPLKSPQRA EADDPQDMACTPSGDSLETKEDQKMSPKATEETGQAQSGQANCQGLSPVSVASKNPOVPSDGGV RLNKSKTDLLVNDNPDPAPLSPELQDFKCNICGYGYYGNDPTDLIKHFRKYHLGLHNRTRQDAEL DSKILALHNMVQFSHSKDFQKVNRSVFSGVLQDINSSRPVLLNGTYDVQVTSGGTFIGIGRKTPDC QGNTKYFRCKFCNFTYMGNSSTELEQHFLQTHPNKIKASLPSSEVAKPSEKNSNKSIPALQSSDSGD LGKWQDKITVKAGDDTPVGYSVPIKPLDSSRQNGTEATSYYWCKFCSFSCESSSSLKLLEHYGKO HGAVQSGGLNPELNDKLSRGSVINQNDLAKSSEGETMTKTDKSSSGAKKKDFSSKGAEDNMVTS YNCQFCDFRYSKSHGPDVIVVGPLLRHYQQLHNIHKCTIKHCPFCPRGLCSPEKHLGEITYPFACRK SNCSHCALLLLHLSPGAAGSSRVKHQCHQCSFTTPDVDVLLFHYESVHESQASDVKQEANHLQGS DGQQSVKESKEHSCTKCDFITQVEEEISRHYRRAHSCYKCRQCSFTAADTQSLLEHFNTVHCQEQD ITTANGEEDGHAISTIKEEPKIDFRVYNLLTPDSKMGEPVSESVVKREKLEEKDGLKEKVWTESSSD DLRNVTWRGADILRGSPSYTQASLGLLTPVSGTQEQTKTLRDSPNVEAAHLARPIYGLAVETKGFL QGAPAGGEKSGALPQQYPASGENKSKDESQSLLRRRRGSGVFCANCLTTKTSLWRKNANGGYVC NACGLYQKLHSTPRPLNIIKQNNGEQIIRRRTRKRLNPEALQAEQLNKQQRGSNEEQVNGSPLERR SEDHLTESHQREIPLPSLSKYEAQGSLTKSHSAQQPVLVSQTLDIHKRMQPLHIQIKSPQESTGDPGN SSSVSEGKGSSERGSPIEKYMRPAKHPNYSPPGSPIEKYQYPLFGLPFVHNDFQSEADWLRFWSKY KLSVPGNPHYLSHVPGLPNPCQNYVPYPTFNLPPHFSAVGSDNDIPLDLAIKHSRPGPTANGASKEK TKAPPNVKNEGPLNVVKTEKVDRSTQDELSTKCVHCGIVFLDEVMYALHMSCHGDSGPFQCSICO HLCTDKYDFTTHIQRGLHRNNAQVEKNGKPKE



breast cancer colon cancer normal tissues

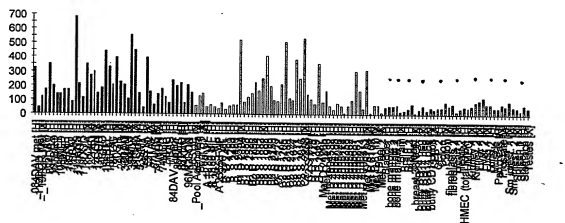
### FIGURE 49A

TAAACTTGGCAAATGACATGCAGGTTCTTCAAGGCAGAATAATTGCAGAAAATCTTCAAAGG TAATTCAAGAACTCCAGAAATCAGGAGACGGAGACATTTTGTCAGTTTTGCAACATTGGACCA AATACA AAGTATTCTTGCTGTGCTCTGGTTTTGGCTGTCCTGGGCACAGAATTGCTGGGA AGCCTCTGTTCGACTGTCAGATCCCCGAGGTTCAGAGGACGGATACAGCAGGAACGAAAAAA CATCCGACCCAACATTATTCTTGTGCTTACCGATGATCAAGATGTGGAGCTGGGGTCCCTGC **AAGTCATGAACAAAACGAGAAAGATTATGGAACATGGGGGGGCCACCTTCATCAATGCC** TTTGTGACTACACCCATGTGCTGCCCGTCACGGTCCTCCATGCTCACCGGGAAGTATGT GCACAÀTCACAATGTCTACACCAACAACGAGAACTGCTCTTCCCCCTCGTGGCAGGCCA TGCATGAGCCTCGGACTTTTGCTGTATATCTTAACAACACTGGCTACAGAACAGCCTTTT TTGGAAAATACCTCAATGAATATAATGGCAGCTACATCCCCCCTGGGTGGCGAGAATGG CTTGGATTAATCAAGAATTCTCGCTTCTATAATTACACTGTTTGTCGCAATGGCATCAAA GAAAAGCATGGATTTGATTATGCAAAGGACTACTTCACAGACTTAATCACTAACGAGAG CATTAATTACTTCAAAATGTCTAAGAGAATGTATCCCCATAGGCCCGTTATGATGGTGAT CAGCCACGCTGCGCCCCACGGCCCCGAGGACTCAGCCCCACAGTTTTCTAAACTGTACC CCAATGCTTCCCAACACATAACTCCTAGTTATAACTATGCACCAAATATGGATAAACACT GGATTATGCAGTACACAGGACCAATGCTGCCCATCCACATGGAATTTACAAACATTCTAC AGCGCAAAAGGCTCCAGÁCTTTGATGTCAGTGGATGATTCTGTGGAGAGGCTGTATAAC ATGCTCGTGGAGACGGGGGAGCTGGAGAATACTTACATCATTTACACCGCCGACCATGG TTACCATATTGGGCAGTTTGGACTGGTCAAGGGGAAATCCATGCCATATGACTTTGATAT TCGTGTGCCTTTTTTTATTCGTGGTCCAAGTGTAGAACCAGGATCAATAGTCCCACAGAT CGTTCTCAACATTGACTTGGCCCCCACGATCCTGGATATTGCTGGGCTCGACACACCTCC TGATGTGGACGCAAGTCTGTCCTCAAACTTCTGGACCCAGAAAAGCCAGGTAACAGGT TTCGAACAACAAGAAGGCCAAAATTTGGCGTGATACATTCCTAGTGGAAAGAGGCAAA TTTCTACGTAAGAAGGAAGAATCCAGCAAGAATATCCAACAGTCAAATCACTTGCCCAAA TATGAACGGGTCAAAGAACTATGCCAGCAGGCCAGGTACCAGACAGCCTGTGAACAACC **GGGGCAGAAGTGGCAATGCATTGAGGATACATCTGGCAAGCTTCGAAATTCACAAGTGTA AAGGACCCAGTGACCTGCTCACAGTCCGGCAGAGCACGCGGAACCTCTACGCTČGCGGC** TTCCATGACAAAGACAAAGAGTGCAGTTGTAGGGAGTCTGGTTACCGTGCCAGCAGAAG CCAAAGAAGAGTCAACGGCAATTCTTGAGAAACCAGGGGACTCCAAAGTACAAGCCCA GATTTGTCCATACTCGGCAGACACGTTCCTTGTCCGTCGAATTTGAAGGTGAAATATATG ACATAAATCTGGAAGAAGAAGAAGAATTGCAAGTGTTGCAACCAAGAAACATTGCTAAG CGTCATGATGAAGGCCACAAGGGGCCAAGAGATCTCCAGGCTTCCAGTGGTGGCAACAG GGGCAGGATGCTGGCAGATAGCAGCAACGCCGTGGGCCCACCTACCACTGTCCGAGTG TCGGCCAGAGCGTGGAAGGACCATAAGGCATACATTGACAAAGAGATTGAAGCTCTGCA **AATGTAGCTGCAGTAAACAAAGCTATTACAATAAAGAGAAAAGGTGTAAAAAAAGCAAGAG AAATTAAAGAGCCATCTTCACCCATTCAAGGAGGCTGCTCAGGAAGTAGATAGCAAACT** CAGAGGAAGGGGAAGAGTGCAGCCTGCCTGGCCTCACTTGCTTCACGCATGACAACAA CCACTGGCAGACAGCCCGTTCTGGAACCTGGGATCTTTCTGTGCTTGCACGAGTTCTA ACAATAACACCTACTGGTGTTTGCGTACAGTTAATGAGACGCATAATTTTCTTTTCTGTG AGTTTGCTACTGGCTTTTTGGAGTATTTTGATATGAATACAGATCCTTATCAGCTCACAA ATACAGTGCACACGGTAGAACGAGGCATTTTGAATCAGCTACACGTACAACTAATGGAG CTCAGAAGCTGTCAAGGATATAAGCAGTGCAACCCAAGACCTAAGAATCTTGATGTTGG GTIMATCAGCCCGTCTCACTGCAGACATCAACTGGCAAGGCCTAGAGGAGCTACACAG TGTGAATGAAAACATCTATGAGTACAGACAAAACTACAGACTTAGTCTGGTGGACTGGA

CTAATTACTTGAAGGATTTAGATAGAGTATTTGCACTGCTGAAGAGTCACTATGAGCAAA ATAAAACAAATAAGACTCAAACTGCTCAAAGTGACGGGTTCTTGGTTGTCTCTGCTGAGC ACGCTGTGTCAATGGAGATGGCCTCTGCTGACTCAGATGAAGACCCAAGGCATAAGGTT GGGAAAACACCTCATTTGACCTTGCCAGCTGACCTTCAAACCCTGCATTTGAACCGACCA ACATTAAGTCCAGAGAGTAAACTTGAATGGAATAACGACATTCCAGAAGTTAATCATTTG **AATTCTGAACACTGGAGAAAAACCGAAAAATGGACGGGGCATGAAGAGACTAATCATCT** GGAAACCGATTTCAGTGGCGATGGCATGACAGAGCTAGAGCTCGGGCCCAGCCCCAGG CTGCAGCCCATTCGCAGGCACCCGAAAGAACTTCCCCAGTATGGTGGTCCTGGAAAGGA CATTTTTGAAGATCAACTATATCTTCCTGTGCATTCCGATGGAATTTCAGTTCATCAGAT GTTCACCATGCCACCGCAGAACACCGAAGTAATTCCAGCATAGCGGGGAAGATGTTGA CCAAGGTGGAGAAGAATCACGAAAAGGAGAAGTCACAGCACCTAGAAGGCAGCGCCTC CTCTTCACTCTCTGATTAGATGAAACTGTTACCTTACCCTAAACACAGTATTTCTTTT TAACTTTTTTATTTGTAAACTAATAAAGGTAATCACAGCCACCAACATTCCAAGCTACCC TGGGTACCTTTGTGCAGTAGAAGCTAGTGAGCATGTGAGCAAGCGGTGTGCACACGGAG TGGCTTGGTTTTGATTTTTTGCTTGTTTGTTTTGTACTAAAACAGTATTATCTTTTG **AATATCGTAGGGACATAAGTATATACATGTTATCCAATCAAGATGGCTAGAATGGTGCCT** GTAATGAAGTTTTGATTCATTTTTAACCACTGGAATTTTTCAATGCCGTCATTTTCAGTTA GATGATTTTGCACTTTGAGATTAAAATGCCATGTCTATTTGATTAGTCTTATTTTTTATT TTTACAGGCTTATCAGTCTCACTGTTGGCTGTCATTGTGACAAAGTCAAATAAACCCCCA AGGACGACACAGTATGGATCACATATTGTTTGACATTAAGCTTTTGCCAGAAAATGTT GCATGTGTTTTACCTCGACTTGCTAAAATCGATTAGCAGAAAGGCATGGCTAATAATGTT GGTGGTGAAAATAAATAAATAAGTAAACAAAAWRAARAWWGCCTGCTCTCTCTGTGCC TAGCCTCAAAGCGTTCATCATACATCATACCTTTAAGATTGCTATATTTTGGGTTATTTTC TTGACAGGAGAAAAAGATCTAAAGATCTTTTATTTTCATCTTTTTTGGTTTTCTTGGCATG **ACTAAGAAGCTTAAATGTTGATAAAATATGACTAGTTTTGAATTTACACCAAGAACTTCT** CAATAAAAGAAATCATGAATGCTCCACAATTTCAACATACCACAAGAGAAGTTAATTTC TTAACATTGTGTTCTATGATTATTTGTAAGACCTTCACCAAGTTCTGATATCTTTTAAAGA CATAGTTCAAAATTGCTTTTGAAAATCTGTATTCTTGAAAATATCCTTGTTGTGTATTAGG TTTTTAAATACCAGCTAAAGGATTACCTCACTGAGTCATCAGTACCCTCCTATTCAGCTC CCCAAGATGATGTTTTTGCTTACCCTAAGAGAGGTTTTCTTCTTATTTTTAGATAATTC **AAGTGCTTAGATAAATTATGTTTTCTTTAAGTGTTTATGGTAAACTCTTTTAAAGAAAATT** TAATATGTTATAGCTGAATCTTTTTGGTAACTTTAAATCTTTATCATAGACTCTGTACATA TTTATGATCATGAATAATGTGCTTTGTAAAAAGATTTCAAGTTATTAGGAAGCATACTCT **GTTTTTTAATCATGTATAATATTCCATGATACTTTTATAGAACAATTCTGGCTTCAGGAAA** AAAAAAAAAAA

## FIGURE 49B

MKYSCCAEMIA ALGIFFIGSI STVRSPRFRGRIQQERKNIRPNIILVLTDDQDVELGSLQVMNKT RKIMEHGGATFINAFVTTPMCCPSRSSMLTGKYVHNHNVYTNNENCSSPSWQAMHEPRTFAVYL NNTGYRTAFFGKYLNEYNGSYIPPGWREWLGLIKNSRFYNYTVCRNGIKEKHGFDYAKDYFTDLI TNESINYFKMSKRMYPHRPVMMVISHAAPHGPEDSAPQFSKLYPNASQHITPSYNYAPNMDKHWI MQYTGPMLPIHMEFTNILQRKRLQTLMSVDDSVERLYNMLVETGELENTYIIYTADHGYHIGQFG LVKGKSMPYDFDIRVPFFIRGPSVEPGSIVPQIVLNIDLAPTILDIAGLDTPPDVDGKSVLKLLDPEKP GNRFRTNKKAKIWRDTFLVERGKFLRKKEESSKNIQQSNHLPKYERVKELCQQARYQTACEQPGQ KWQCIEDTSGKLRIHKCKGPSDLLTVRQSTRNLYARGFHDKDKECSCRESGYRASRSQRKSQRQF LRNQGTPKYKPRFVHTRQTRSLSVEFEGEIYDINLEEEEELQVLQPRNIAKRHDEGHKGPRDLQASS GGNRGRMLADSSNAVGPPTTVRVTHKCFILPNDSIHCERELYQSARAWKDHKAYIDKEIEALQDKI KNLREVRGHLKRRKPEECSCSKQSYYNKEKGVKKQEKLKSHLHPFKEAAQEVDSKLQLFKENNR RRKKERKEKRRQRKGEECSLPGLTCFTHDNNHWQTAPFWNLGSFCACTSSNNNTYWCLRTVNET HNFLFCEFATGFLEYFDMNTDPYQLTNTVHTVERGILNQLHVQLMELRSCQGYKQCNPRPKNLDV GNKDGGSYDLHRGQLWDGWEG



# breast cancer colon cancer normal tissues

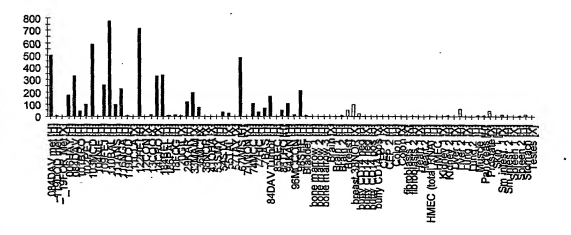
AACCGAGAAGCGCTCCGTAAAGCCATCCGCACGCTCAGAAAGGCCGTCCACAGGGAGCAGTTTCACCTCCAGCTCTCAGG CATGAACCTCGACGTGGCTAAAAAGCCTCCCAGAACATCTGAACGCCAGGCAGAGTCCTGTGGAGTGGGCCAGGGTCATG CAGAAAACCAATGTGTCAGTTGCAGGGCTGGGACCTATTATGATGGAGCACGAGAACGCTGCATTTTATGTCCAAATGGA ACCTTCCAAAATGAGGAAGGACAAATGACTTGTGAACCATGCCCAAGACCAGGAAATTCTGGGGCCCTGAAGACCCCAGA AGCTTGGAATATGTCTGAATGTGGAGGKCTGTGTCAACCTACTGAATATTCTGCAGATGGCTTTGCACCTTGCCAGCTCT GTGCCCTGGGCASGTTCCAGCCTGAAGCTGGTCGAACTTCCTGCTTCCCCTGTGGAGGAGGCCTTGCCACCAAACATCAG GGAGCTACTTCCTTTCAGGACTGTGAAACCAGAGTTCAATGTTCACCTGGACATTTCTACAACACCACCACTCACCGATG TATTCGTTGCCCAGTGGGAACATACCAGCCTGAATTTGGAAAAAATAATTGTGTTTCTTGCCCAGGAAATACTACGACTG CCTGATCGTGGTCCCTGAGATCTTCCTGCCCATAGAGGACGACTGTGGGGGACTATCTGGTGATGCGGAAAACCTCTTCAT CCAATTCTGTGACAACATATGAAACCTGCCAGACCTACGAACGCCCCATCGCCTTCACCTCCAGGTCAAAGAAGCTGTGG ATTCAGTTCAAGTCCAATGAAGGGAACAGCGCTAGAGGGTTCCAGGTCCCATACGTGACATATGATGAGGACTACCAGGA ACTCATTGAAGACATAGTTCGAGATGGCAGGCTCTATGCATCTGAGAACCATCAGGAAATACTTAAGGATAAGAAACTTA TCAAGGCTCTGTTTGATGTCCTGGCCCATCCCCAGAACTATTTCAAGTACACAGCCCAGGAGTCCCGAGAGATGTTTCCA AGATCGTTCATCCGATTGCTACGTTCCAAAGTGTCCAGGTTTTTGAGACCTTACAAATGACTCAGCCCACGTGCCACTCA AGCATCGTGGATGTAGACTGAGAATGGCTTTGAGTGGCATCAGCTTCTCACTGCTGTGGGCGGATGTCTTGGATAGATCA AGGGCTGGCTGAGCTTGGTCAGCCTAGGTGAGACTCACCTGTCCTTCTGGGGTCTTACTCCTCCTCAAGGAGTC TGTAGTGGAAAGGAGGCCACAGAATAAGCTGCTTATTCTGAAACTTCAGCTTCCTCTAGCCCGGCCCTCTCTAAGGGAGC CCACCTTGAGACCTGGGAGGACTCAGTTTCTCCACAGCCTTCTCCAGCCTGTGTGATACAAGTTTGATCCCAGGAACTTG AA

### FIGURE 5 2

### FIGURE 54

...MNLDVAKKPPRTSERQAESCGVGQGHAENQCVSCRAGTYYDGARERCILCPNGTFQNEEGQMTCEPCPRPGNSGALK TPEAWNMSECGGLCQPTEYSADGFAPCQLCALGXFQPEAGRTSCFPCGGGLATKHQGATSFQDCETRVQCSPGHFYNTTT HRCIRCPVGTYQPEFGKNNCVSCPGNTTTDFDGSTNITQCKNRRCGGELGDFTGYIESPNYPGNYPANTECTWTINPPPK RRILIVVPEIFLPIEDDCGDYLVMRKTSSSNSVTTYETCQTYERPIAFTSRSKKLWIQFKSNEGNSARGFQVPYVTYDED YQELIEDIVRDGRLYASENHQEILKDKKLIKALFDVLAHPQNYFKYTAQESREMFPRSFIRLLRSKVSRFLRPYK.

human_BCO2 mouse_BCO2	MNLDVAKKPPRTSERQAESCGVGQGHAENQCVSCRAGTYYDGARERCILCPNGTFQNEEG
human_BCO2 mouse_BCO2	QMTCEPCPRPGNSGALKTPEAWNMSECGGLCQPTEYSADGFAPCQLCALGXFQPEAGRTS
human_BCO2 mouse_BCO2	CFPCGGGLATKHQGATSFQDCETRVQCSPGHFYNTTTHRCIRCPVGTYQPEFGKNNCVSC
human_BCO2 mouse_BCO2	PGNTTTDFDGSTNITQCKNRRCGGELGDFTGYIESPNYPGNYPANTECTWTINPPPKRRI
human_BCO2 mouse_BCO2	LIVVPEIFLPIEDDCGDYLVMRKTSSSNSVTTYETCQTYERPIAFTSRSKKLWIQFKSNE LIVVPEIFLPIEDDCGDYLVMRKTSSSNSVTTYETCQTYERPIAFTSRSKKLWIQFKSNE ************************************
human_BCO2 mouse_BCO2	GNSARGFQVPYVTYDEDYQELIEDIVRDGRLYASENHQEILKDKKLIKALFDVLAHPQNY GNSARGFQVPYVTYDEDYQELIEDIVRDGRLYASENHQEILKDKKLIKALFDVLAHPQNY ************************************
human_BCO2 mouse_BCO2	FKYTAQESREMFPRSFIRLLRSKVSRFLRPYK FKYTAQESREMFPRSFIRLLRSKVSRFLRPYK ************************************



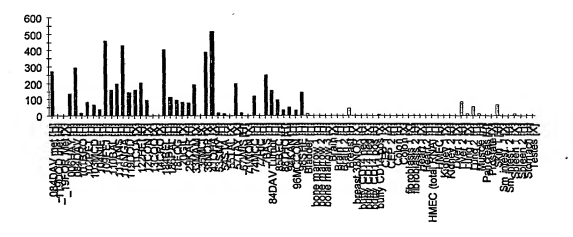
breast cancer normal tissues

CAGCGCCGCTGAATTCTAGGGCGGGTTCGCGCCCCGAAGGCTGAGAGCTGGCGCTGCTCGTGCCCTGTG TGCCAGACGGCGGAGCTCCGCGGCCGGGCCCCGGCTTTGCTGCCGACTGGAGTTTGGGGGAAG AAACTCTCCTGCGCCCCAGAAGATTTCTTCCTCGGCGAAGGGACAGCGAAAGATGAGGGTGGCAGGAAGA GAAGGCGCTTTCTGTCTGCCGGGGTCGCAGCGCGAGAGGGCAGTGCCATGTTCCTCCCATCCTAGTGGC TGCCGGCACATGCCCTGGAACATCACGCGGATGCCCAACCACCTGCACCACAGCACGCAGGAGAACGCCA TCCTGGCCATCGAGCAGTACGAGGAGCTGGTGGACGTGAACTGCAGCGCCGTGCTGCGCGTTCTTCTTCTG TGCCATGTACGCGCCCATTTGCACCCTGGAGTTCCTGCACGACCCTATCAAGCCGTGCAAGTCGGTGTGC CAACGCGCGCGCGACGACTGCGAGCCCCTCATGAAGATGTACAACCACAGCTGGCCCGAAAGCCTGGCCT GCGACGAGCTGCCTGTCTATGACCGTGGCGTGTGCATTTCGCCTGAAGCCATCGTCACGGACCTCCCGGA GGATGTTAAGTGGATAGACATCACCAGACATGATGGTACAGGAAAGGCCTCTTGATGTTGACTGTAAA CGCCTAAGCCCCGATCGGTGCAAGTGTAAAAAGGTGAAGCCAACTTTGGCAACGTATCTCAGCAAAAACT ACAGCTATGTTATTCATGCCAAAATAAAAGCTGTGCAGAGGAGTGGCTGCAATGAGGTCACAACGGTGGT GGATGTAAAAGAGATCTTCAAGTCCTCATCACCCATCCCTCGAACTCAAGTCCCGCTCATTACAAATTCT TCTTGCCAGTGTCCACACATCCTGCCCCATCAAGATGTTCTCATCATGTGTTACGAGTGGCGTTCAAGGA TGATGCTTCTTGAAAATTGCTTAGTTGAAAAATGGAGAGATCAGCTTAGTAAAAGATCCATACAGTGGGA AGAGAGGCTGCAGGAACAGCGGAGAACAGTTCAGGACAAGAAGAAAACAGCCGGGCGCACCAGTCGTAGT GGAGTGCCCAGAAGAGAACAAACCCGAAAAGAGTGTGAGCTAACTAGTTTCCAAAGCGGAGACTTCCGAC TTCCTTACAGGATGAGGCTGGGCATTGCCTGGGACAGCCTATGTAAGGCCATGTGCCCCTTGCCCTAACA ACTCACTGCAGTGCTCTTCATAGACACATCTTGCAGCATTTTTCTTAAGGCTATGCTTCAGTTTTTCTTT GTAAGCCATCACAAGCCATAGTGGTAGGTTTGCCCTTTGGTACAGAAGGTGAGTTAAAGCTGGTGGAAAA GGCTTATTGCATTGCATTCAGAGTAACCTGTGTGCATACTCTAGAAGAGTAGGGAAAATAATGCTTGTTA TATGTTTTATTACCTTTTGATATCTGTTGTTGCAATGTTAGTGATGTTTTAAAATGTGATGAAAATATAA TGTTTTTAAGAAGGAACAGTAGTGGAATGAATGTTAAAAGATCTTTATGTGTTTATGGTCTGCAGAAGGA TTTTTGTGATGAAAGGGGATTTTTTGAAAAATTAGAGAAGTAGCATATGGAAAATTATAATGTGTTTTTT AAAAAGGAGAGGCAGACAATGTCTGGATTCCTGTTTTTTGGTTACCTGATTTCCATGATCATGATGCTTC TTGTCAACACCCTCTTAAGCAGCACCAGAAACAGTGAGTTTGTCTGTACCATTAGGAGTTAGGTACTAAT TAGTTGGCTAATGCTCAAGTATTTTATACCCACAAGAGAGGTATGTCACTCATCTTACTTCCCAGGACAT CCACCCTGAGAATAATTTGACAAGCTTAAAAATGGCCTTCATGTGAGTGCCAAATTTTGTTTTTCTTCAT TTAAATATTTCTTTGCCTAAATACATGTGAGAGGAGTTAAATATAAATGTACAGAGAGGAAAGTTGAGT TTTGCTAACACAGTAAGCATGTATTTTATAAGGCATTCAATAAATGCACAACGCCCAAAGGAAATAAAAT CCTATCTAATCCTACTCTCCACTACACAGAGGTAATCACTATTAGTATTTTTGGCATATTATTCTCCAGGT GTTTGCTTATGCACTTATAAAATGATTTGAACAAATAAAACTAGGAACCTGTATACATGTGTTTCATAAC CTGCCTCCTTTGCTTGGCCCTTTATTGAGATAAGTTTTCCTGTCAAGAAAGCAGAAACCATCTCATTTCT AACAGCTGTGTTATATTCCATAGTATGCATTACTCAACAAACTGTTGTGCTATTGGATACTTAGGTGGTT TCTTCACTGACAATACTGAATAAACATCTCACCGGAATTC

ATGTTCCTCTCCATCCTAGTGGC

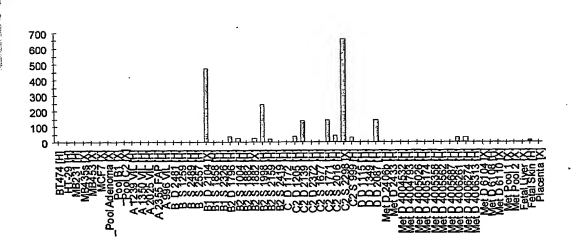
## FIGURE 5.8

MFLSILVALCLWLHLALGVRGAPCEAVRIPMCRHMPWNITRMPNHLHHSTQENAILAIEQYEELVDVNCS AVLRFFFCAMYAPICTLEFLHDPIKPCKSVCQRARDDCEPLMKMYNHSWPESLACDELPVYDRGVCISPE AIVTDLPEDVKWIDITPDMMVQERPLDVDCKRLSPDRCKCKKVKPTLATYLSKNYSYVIHAKIKAVQRSG CNEVTTVVDVKEIFKSSSPIPRTQVPLITNSSCQCPHILPHQDVLIMCYEWRSRMMLLENCLVEKWRDQL SKRSIQWEERLQEQRRTVQDKKKTAGRTSRSNPPKPKGKPPAPKPASPKKNIKTRSAQKRTNPKRV



### breast cancer normal tissues

## FIGURE 60

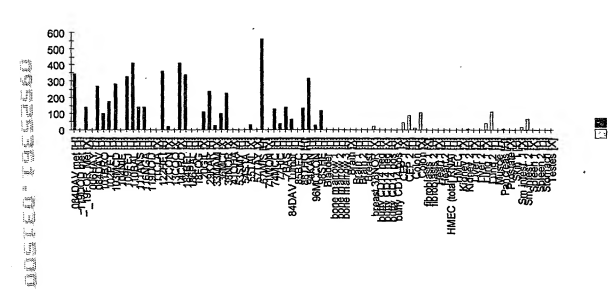


cell lines
colon cancer
fetal tissues

FIGURE 61

## FIGURE 63

MMLHSALGLCLLLVTVSSNLAIAIKKEKRPPQTLSRGWGDDITWVQTYEEGLFYAQKSKKPLMVIHHLEDCQYSQALKKV FAQNEEIQEMAQNKFIMLNLMHETTDKNLSPDGQYVPRIMFVDPSLTVRADIAGRYSNRLYTYEPRDLPLLIENMKKALR LIQSEL

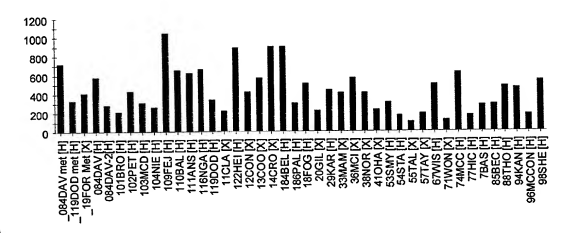


breast cancer normal tissues

GGCGGCGCGCTCCCCGGGAGGGCCCGGCGGGAACGGCCGCGATCGGGGCCGCAGCCACG CTGCCGAGGCGCCCGGGGACCCGCAGGCGGCCGCTCTTGCTGGCCCCTATGGACGTGGGG GAGGAGCCGCTGGAGAAGGCGGCGCGCCCCCACTGCCAAGGACCCCAACACCTATAAAGT **ACTCTCGCTGGTATTGTCAGTATGTGTTTAACAACAATACTTGGTTGTATATTTTGGGTTGAAA** CCAAGCTGTGCCAAAGAAGTTAAAAGTTGCAAAGGTCGCTGTTTCGAGAGAACATTTGGGAA CTGTCGCTGTGATGCTGCCTGTGTTGAGCTTGGAAACTGCTGTTTAGATTACCAGGAGACGTG CATAGAACCAGAACATATATGGACTTGCAACAAATTCAGGTGTGGTGAGAAAAGGTTGACCA GAAGCCTCTGTGCCTGTTCAGATGACTGCAAGGACAAGGGCGACTGCTGCATCAACTACAGTT CTGTGTGTCAAGGTGAGAAAAGTTGGGTAGAAGAACCATGTGAGAGCATTAATGAGCCACAG <u>ATTTACACACTTGGGGTGGACTTCTTCCTGTTATTAGCAAACTAAAAAAATGTGGAACATATA</u> CTAAAAACATGAGACCGGTATATCCAACAAAAACTTTCCCCAATCACTACAGCATTGTCACCG CCTTTTCACTTAAAAGTAAAGAGAAATTTAATCCTGAGTGGTACAAAGGAGAACCAATTTGGG TCACAGCTAAGTATCAAGGCCTCAAGTCTGGCACATTTTTCTGGCCAGGATCAGATGTGGAAA TTAACGGAATTTTCCCAGACATCTATAAAATGTATAATGGTTCAGTACCATTTGAAGAAAGGA TTTTAGCTGTTCTTCAGTGGCTACAGCTTCCTAAAGATGAAAGACCACACTTTTACACTCTGTA TTTAGAAGAACCAGATTCTTCAGGTCATTCATATGGACCAGTCAGCAGTGAAGTCATCAAAGC CTTGCAGAGGGTTGATGGTATGGTATGCTGATGGATGGTCTGAAAGAGCTGAACTTGCA CAGATGCCTGAACCTCATCCTTATTTCAGATCATGGCATGGAACAAGGCAGTTGTAAGAAATA CATATATCTGAATAAATATTTGGGGGATGTTAAAAATATTAAAGTTATCTATGGACCTGCAGC TCGATTGAGACCCTCTGATGTCCCAGATAAATACTATTCATTTAACTATGAAGGCATTGCCCG AAATCTTTCTTGCCGGGAACCAAACCAGCACTTCAAACCTTACCTGAAACATTTCTTACCTAA GCGTTTGCACTTTGCTAAGAGTGATAGAATTGAGCCCTTGACATTCTATTTGGACCCTCAGTGG CAACTTGCATTGAATCCCTCAGAAAGGAAATATTGTGGAAGTGGATTTCATGGCTCTGACAAT GTATTTTCAAATATGCAAGCCCTCTTTGTTGGCTATGGACCTGGATTCAAGCATGGCATTGAG GCTGACACCTTTGAAAACATTGAAGTCTATAACTTAATGTGTGATTTACTGAATTTGACACCG GCTCCTAATAACGGAACTCATGGAAGTCTTAACCACCTTCTAAAGAATCCTGTTTATACGCCA AAGCATCCCAAAGAAGTGCACCCCTGGTACAGTGCCCCTTCACAAGAAACCCCAGAGATAA CCTTGGCTGCTCATGTAACCCTTCGATTTTGCCGATTGAGGATTTTCAAACACAGTTCAATCTG ACTGTGGCAGAAGAGAGATTATTAAGCATGAAACTTTACCCTATGGAAGACCTAGAGTTCTC CAGAAGGAAAACACCATCTGTCTTCTTTCCCAGCACCAGTTTATGAGTGGATACAGCCAAGAC ATCITAATGCCCCTTTGGACATCCTATACCGTGGACAGAATGACAGTTTCTCTACGGAAGAC TTCTCCAACTGTCTGTACCAGGACTTTAGAATTCCTCTTAGTCCTGTCCATAAATGTTCATTTTA TGGAATATATTCTGAAGCTTTGCTTACTACAAATATAGTGCCAATGTACCAGAGTTTTCAAGTT GTCGTCAGTGGTCCTGTGTTTGACTTTGATTATGATGGACGTTGTGATTCCTTAGAGAATCTGA GGCAAAAAAGAAGAGTCATCCGTAACCAAGAAATTTTGATTCCAACTCACTTCTTTATTGTGC TAACAAGCTGTAAAGATACATCTCAGACGCCTTTGCACTGTGAAAACCTAGACACCTTAGCTT TCATTTTGCCTCACAGGACTGATAACAGCGAGAGCTGTGTGCATGGGAAGCATGACTCCTCAT GGGTTGAAGAATTGTTAATGTTACACAGAGCACGGATCACAGATGTTGAGCACATCACTGGA CTCAGCTTCTATCAACAAAGAAAAGAGCCAGTTTCAGACATTTTAAAGTTGAAAACACATTTG CCAACCTTTAGCCAAGAAGAC TATGTTTTTTATCCCCAAACACCATGAATCTTTTTGAGA GAACCTTATATTTTATAGTCCTCTAGCTACACTATTGCATTGTTCAGAAACTGTCGA:CCAG AGT:TAGAACGGAGCCCTCGGTGATGCGGACATCTCAGGGAAACTTGCGTACTCAGCACAGCA GTGGAGAGTGTTCCTGTTGAATCTTGCACATATTTGAATGTGTAAGCATTGTATACATTGATCA AGTTCGGGGGAATAAAGACAGACCACACCTAAAACTGCCTTTCTGCTTCTCTTAAAGGAGAAG ATGGCATTGGGCAAACAGTAGACTTATAGTAGGGTTGGGGTAGCCCATGTTATGTGACTATCT TTATGAGAATTTTAAAGTGGTTCTGGATATCTTTTAACTTGGAGTTTCATTTCTTTTCATTGTAA TCAAAAAAAAATTAACAGAAGCCAAAATACTTCTGAGACCTTGTTTCAATCTTTGCTGTATA TCCCCTCAAAATCCAAGTTATTAATCTTATGTGTTTTTCTTTTTAATTTTTTGATTGGATTTCTTT AGATTTAATGGTTCAAATGAGTTCAACTTTGAGGGACGATCTTTGAATATACTTACCTATTATA AAATCTTACTTTGTATTTGTATTT

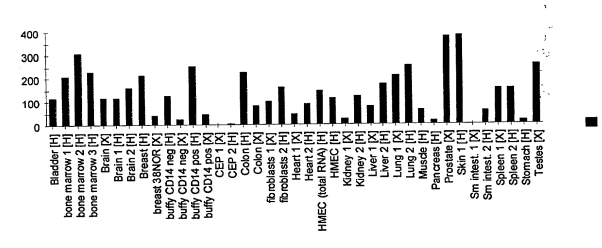
MERDGCAGGGSRGGEGGRAPREGPAGNGRDRGRSHAAEAPGDPQAAASLLAPMDVGEEPLEKA ARARTAKDPNTYKVLSLVLSVCVLTTILGCIFGLKPSCAKEVKSCKGRCFERTFGNCRCDAACVEL GNCCLDYQETCIEPEHIWTCNKFRCGEKRLTRSLCACSDDCKDKGDCCINYSSVCQGEKSWVEEP CESINEPQCPAGFETPPTLLFSLDGFRAEYLHTWGGLLPVISKLKKCGTYTKNMRPVYPTKTFPNH YSIVTGLYPESHGIIDNKMYDPKMNASFSLKSKEKFNPEWYKGEPIWVTAKYQGLKSGTFFWPGS DVEINGIFPDIYKMYNGSVPFEERILAVLQWLQLPKDERPHFYTLYLEEPDSSGHSYGPVSSEVIKA LQRVDGMVGMLMDGLKELNLHRCLNLILISDHGMEQGSCKKYIYLNKYLGDVKNIKVIYGPAAR LRPSDVPDKYYSFNYEGIARNLSCREPNQHFKPYLKHFLPKRLHFAKSDRIEPLTFYLDPQWQLAL NPSERKYCGSGFHGSDNVFSNMQALFVGYGPGFKHGIEADTFENIEVYNLMCDLLNLTPAPNNGT HGSLNHLLKNPVYTPKHPKEVHPLVQCPFTRNPRDNLGCSCNPSILPIEDFQTQFNLTVAEEKIIKHE TLPYGRPRVLQKENTICLLSQHQFMSGYSQDILMPLWTSYTVDRNDSFSTEDFSNCLYQDFRIPLSP VHKCSFYKNNTKVSYGFLSPPQLNKNSSGIYSEALLTTNIVPMYQSFQVIWRYFHDTLLRKYAEER NGVNVVSGPVFDFDYDGRCDSLENLRQKRRVIRNQEILIPTHFFIVLTSCKDTSQTPLHCENLDTLA FILPHRTDNSESCVHGKHDSSWVEELLMLHRARITDVEHITGLSFYQQRKEPVSDILKLKTHLPTFS QED.

GGTTTTCAA GGAACATTTTGATGCATCACTTAGTACCTATTTCAAGGCATTGCTAGGCCCTC GAGATACTAGAGTAAAAGGATGGTTTCTTCTGGACAATTATATACCCACATTTATCTGCTCTGT CATATATTTACTAATTGTATGGCTGGGACCAAAATACATGAGGAATAAACAGCCATTCTCTTG CCGGGGGATTTTAGTGGTGTATAACCTTGGACTCACACTGCTGTCTCTGTATATGTTCTGTGAG TTAGTAACAGGAGTATGGGAAGGCAAATACAACTTCTTCTGTCAGGGCACACGCACCGCAGG AGAATCAGATATGAAGATTATCCGTGTCCTCTGGTGGTACTACTTCTCCAAACTCATAGAATTT ATGGACACTTTCTTCATCCTGCGCAAGAACAACCACCAGATCACGGTCCTGCACGTCTAC CACCATGCCTCGGTGCTGAACATCTGGTGGTTTGTGATGAACTGGGTCCCCTGCGGCCACTCTT ATTTTGGTGCCACACTTAATAGCTTCATCCACGTCCTCATGTACTCTTACTATGGTTTGTCGTC AGTCCCTTCCATGCGTCCATACCTCTGGTGGAAGAAGTACATCACTCAGGGGCAGCTGCTTCA GTTTGTGCTGACAATCATCCAGACCAGCTGCGGGGTCATCTGGCCGTGCACATTCCCTCTTGGT TGGTTGTATTTCCAGATTGGATACATGATTTCCCTGATTGCTCTCTCACAAACTTCTACATTC AGACCTACAACAAGAAAGGGGCCTCCCGAAGGAAAGACCACCTGAAGGACCACCAGAATGG GTCCATGGCTGTGAATGGACACACCAACAGCTTTTCACCCCTGGAAAACAATGTGAAGCC AAGGAAGCTGCGGAAGGAT AGTCAAAGAATTGAAACCCTCCAAACCACGTCATCTGATT GTAAGCACAATATGAGTTGTGCCCCAATGCTCGTTAACAGCTGCTGTAACTAGTCTGGCCTAC AATAGTGTGATTCAAAGGGCGAATTCTTTCATCAATTCAAAACCCCTAGAAAACGTATACAGA TTATATAAGTAGGGATAAGATTTCTAACATTTCTGGGCTCTCTGACCCCTGCGCTAGACTGTGG AAAGGGAGTATTATTATAGTATACAACACTGCTGTTGCCTTATTAGTTATAACATGATAGGTG GCATGTGATTGTAAATGTAAATTTGTACAATGTTGTTATGGTAGAGAAACACACATGCCTTAA AATTTAAAAAGCAGGCCCAAAGCTTATTAGTTTAAATTAGGGTATGTTTCAAGTTTGTATTA ATTTGTAATAGCTCTGTTTAGAAAAAATCAAAGACCATGATTTATGAAACTAATGTGACATAA TTTCCAGTGACTTGTTGATGTGAAATCAGACACGGCACCTTCAGTTTTGTACTATTGGCTTTGA ATCAAGCAGGCTCAAATCTAGTGGAACAGTCAGTTTAACTTTTTAACAGATCTTATTTTTTAT TTTGAGTGCCACTATTAATGTAAAAAGGGGGGGGCTCTACAGCAGTCGTGATGAAACTTAAAT ATATATTCTTTGTCCTCGAGATTTTAGGAAGGGTGTAGGGTGAGTAGGCCATTTTTAATTTCTG AAGTGCTAAGTGTTTTTATACAGCAAACAAAAAGTCAATTTTGCTTTCCACCAGTGCGAGAGA GGATGTATACTTTTCAAGAGAGATGATTGCCTATTTACCGTTTGACAGAGTCCCGTAGATGAG ACAGATTTCCTTGTAAAATGTGCCTAGTTTACCAAAATTAACAAAGGGGGGGAAAGGACCTTA GAACTTTTTAAGGTAAAATCAAATATAGCTACAGCATAAGAGAATCGAGAAATTTGATAGAG GTAACTTGTTTAATGTAAATCTAATAGTACTTGTAATTTCTTCTGCTTAGAATCTAAAGATGT GTTTAGAACCTCTTGTTTAAAAATAATAGACTGCTTATCATAAAATCACATCTCACACATTTGA GGCAGTGGTCAAACAGGTAAAGCCTATGATGTGTGTCATTTTAAAGTGTCGGAATTTAGCCTC TGAATACCTTCTCCATTGGGGGAAAGATATTCTTGGAACCACTCATGACATATCTTAGAAGGT CATTGACAATGTATAAACTAATTGTTGGTTTGATATTTATGTAAATATCAGTTTACCATGCTTT AATTTTGCACATTCGTACTATAGGGAGCCTATTGGTTCTCTATTAGTCTTGTGGGTTTTCTGTTT GAAAAGGAGTCATGGCATCTGTTTACATTTACCTTATCAAACCTAGAATGTGTATATTTATAA ATGTATGTCTTCATTGCTAGGTACTAATTTGCAGATGTCTTTACATATTTCAATACAGAAACTA TAACATTCAATAGTGTGCTGTCAAAGTGTGCTTAGCTCACCTGGATATACCTACATTGTTAAAT 



breast cancer

## FIGURE 70



normal tissues

## FIGURE 72

AGAGATGGGGGTCTCACTATGTTGCCCAGGCTGGTCTCÄAACTCCTGGGCTCAAGCGATCCTTTGGCCTCG GTTTTCTAATTAGAAAAGTAATATCTACTCATTGTAAAACTCAAACAGTGCAGAAATGTAGAAAGTAGAA AGTGTAAGTCCCTGGTTGTCCCTTCTGCCTGAGCACAAGCACTGCTCACAGTTTGATGTATATCCTTCCAG AGACTCTCAAATTTAAGCAAATAATTTTTATTACCATGTCTTTTTATTTGAAGACGTACATTTGCCTCCAA AGTTCAACACAAGTTCAACTGACCATATCCTTCCATGACCTGAATAGATGCTATCCTTTATCACGATGTTC ANTTGCCTTTGANAGAGAGTAGTCCAGGTATATTCCTGATCAAAATTTGGCATTTTTGATGATACTACTCT CATTCAAATTGCAAATGGAAACTTGAAACCCACAATCTAATGAGGAATGTACTGGAAAAATAATCTGAAGA GTTGACAAATTGTGTACTAGATTGAACACATGGAATGCAATGCCAATGAGACTTTCTGCACTAAAACTTAT ACGCCTGAAGGATATTACTAGTGCCTAATATTGAGTATGAGTCACTGCGTGTTCGCATCAACTTGGAAGTG CAGTANTIGTTATANNATTAATCAGTGCAGCCAACATTATTATGAATCACATCTTTGAAACTGTGCAGTA GCN1'ATACATATATATTTTAAATAACATTTTTCACAGATTTCCAGAGTTACTGTTGAAATCTGCA'I'CACC AAAAAAAAAAAAGCAAGATTTTTTTAACAATGTAGÀÇACTCTTCAGACCCAGTAATCTGCGTGTGATTT CCTATTTGTACATTCCCAAGAGACTTTAGCAGTCACCAGCCTTAATGCATGTACAGGATATTATTGTGACT TAATTTATCTGCAGTTTTTAATCCATGTGAAATTGGGAATTTTTAACCGAACTTGGATTAACCATGCCTGC